In the Senate of the United States,

October 14, 1999.

Resolved. That the bill from the House of Representatives (H.R. 2990) entitled "An Act to amend the Internal Revenue Code of 1986 to allow individuals greater access to health insurance through a health care tax deduction, a longterm care deduction, and other health-related tax incentives, to amend the Employee Retirement Income Security Act of 1974 to provide access to and choice in health care through association health plans, to amend the Public Health Service Act to create new pooling opportunities for small employers greater access to health coverage through obtain to HealthMarts; to amend title I of the Employee Retirement Income Security Act of 1974, title XXVII of the Public Health Service Act, and the Internal Revenue Code of 1986 to protect consumers in managed care plans and other health coverage; and for other purposes.", do pass with the following

AMENDMENT:

Strike out all after the enacting clause and insert:

1 SECTION 1. SHORT TITLE; TABLE OF CONTENTS.

- 2 (a) SHORT TITLE.—This Act may be cited as the "Pa-
- 3 tients' Bill of Rights Plus Act".
- 4 (b) TABLE OF CONTENTS.—The table of contents for
- 5 this Act is as follows:

Sec. 1. Short title; table of contents.

TITLE I—PATIENTS' BILL OF RIGHTS

Subtitle A—Right to Advice and Care

Sec. 101. Patient right to medical advice and care.

"SUBPART C-PATIENT RIGHT TO MEDICAL ADVICE AND CARE

- "Sec. 721. Patient access to emergency medical care.
- "Sec. 722. Offering of choice of coverage options.
- "Sec. 723. Patient access to obstetric and gynecological care.
- "Sec. 724. Patient access to pediatric care.
- "Sec. 725. Timely access to specialists.
- "Sec. 726. Continuity of care.
- "Sec. 727. Protection of patient-provider communications.
- "Sec. 728. Patient's right to prescription drugs.
- "Sec. 729. Self-payment for behavioral health care services.
- "Sec. 730. Coverage for individuals participating in approved cancer clinical trials.
- "Sec. 730A. Prohibiting discrimination against providers.
- "Sec. 730B. Generally applicable provision.".
- Sec. 102. Conforming amendment to the Internal Revenue Code of 1986.

"SUBCHAPTER C-PATIENT RIGHT TO MEDICAL ADVICE AND CARE

- "Sec. 9821. Patient access to emergency medical care.
- "Sec. 9822. Offering of choice of coverage options.
- "Sec. 9823. Patient access to obstetric and gynecological care.
- "Sec. 9824. Patient access to pediatric care.
- "Sec. 9825. Timely access to specialists.
- "Sec. 9826. Continuity of care.
- "Sec. 9827. Protection of patient-provider communications.
- "Sec. 9828. Patient's right to prescription drugs.
- "Sec. 9829. Self-payment for behavioral health care services.
- "Sec. 9830. Coverage for individuals participating in approved cancer clinical trials.
- "Sec. 9830A. Prohibiting discrimination against providers.
- "Sec. 9830B. Generally applicable provision.".
- Sec. 103. Effective date and related rules.

Subtitle B-Right to Information About Plans and Providers

- Sec. 111. Information about plans.
- Sec. 112. Information about providers.

Subtitle C-Right to Hold Health Plans Accountable

Sec. 121. Amendment to Employee Retirement Income Security Act of 1974.

TITLE II—WOMEN'S HEALTH AND CANCER RIGHTS

Sec. 201. Women's health and cancer rights.

TITLE III—GENETIC INFORMATION AND SERVICES

- Sec. 301. Short title.
- Sec. 302. Amendments to Employee Retirement Income Security Act of 1974.

- Sec. 303. Amendments to the Public Health Service Act.
- Sec. 304. Amendments to the Internal Revenue Code of 1986.

TITLE IV—HEALTHCARE RESEARCH AND QUALITY

Sec. 401. Short title.

Sec. 402. Amendment to the Public Health Service Act.

"TITLE IX—AGENCY FOR HEALTHCARE RESEARCH AND QUALITY

"PART A-ESTABLISHMENT AND GENERAL DUTIES

- "Sec. 901. Mission and duties.
- "Sec. 902. General authorities.

"Part B—Healthcare Improvement Research

- "Sec. 911. Healthcare outcome improvement research.
- "Sec. 912. Private-public partnerships to improve organization and delivery.
- "Sec. 913. Information on quality and cost of care.
- "Sec. 914. Information systems for healthcare improvement.
- "Sec. 915. Research supporting primary care and access in underserved areas.
- "Sec. 916. Clinical practice and technology innovation.
- "Sec. 917. Coordination of Federal government quality improvement efforts.

"PART C—GENERAL PROVISIONS

- "Sec. 921. Advisory Council for Healthcare Research and Quality.
- "Sec. 922. Peer review with respect to grants and contracts.
- "Sec. 923. Certain provisions with respect to development, collection, and dissemination of data.
- "Sec. 924. Dissemination of information.
- "Sec. 925. Additional provisions with respect to grants and contracts.
- "Sec. 926. Certain administrative authorities.
- "Sec. 927. Funding.
- "Sec. 928. Definitions.".
- Sec. 403. References.

TITLE V-ENHANCED ACCESS TO HEALTH INSURANCE COVERAGE

- Sec. 501. Full deduction of health insurance costs for self-employed individuals.
- Sec. 502. Full availability of medical savings accounts.
- Sec. 503. Permitting contribution towards medical savings account through Federal employees health benefits program (FEHBP).
- Sec. 504. Carryover of unused benefits from cafeteria plans, flexible spending arrangements, and health flexible spending accounts.

TITLE VI—PROVISIONS RELATING TO LONG-TERM CARE INSURANCE

- Sec. 601. Inclusion of qualified long-term care insurance contracts in cafeteria plans, flexible spending arrangements, and health flexible spending accounts.
- Sec. 602. Deduction for premiums for long-term care insurance.
- Sec. 603. Study of long-term care needs in the 21st century.

TITLE VII—INDIVIDUAL RETIREMENT PLANS

Sec. 701. Modification of income limits on contributions and rollovers to Roth IRAs.

TITLE VIII—REVENUE PROVISIONS

- Sec. 801. Modification to foreign tax credit carryback and carryover periods.
- Sec. 802. Limitation on use of non-accrual experience method of accounting.
- Sec. 803. Returns relating to cancellations of indebtedness by organizations lending money.
- Sec. 804. Extension of Internal Revenue Service user fees.
- Sec. 805. Property subject to a liability treated in same manner as assumption of liability.
- Sec. 806. Charitable split-dollar life insurance, annuity, and endowment contracts.
- Sec. 807. Transfer of excess defined benefit plan assets for retiree health benefits.
- Sec. 808. Limitations on welfare benefit funds of 10 or more employer plans.
- Sec. 809. Modification of installment method and repeal of installment method for accrual method taxpayers.
- Sec. 810. Inclusion of certain vaccines against streptococcus pneumoniae to list of taxable vaccines.

TITLE IX—MISCELLANEOUS PROVISIONS

Sec. 901. Medicare competitive pricing demonstration project.

1	TITLE I—PATIENTS' BILL OF
2	RIGHTS
3	Subtitle A—Right to Advice and
4	Care
5	SEC. 101. PATIENT RIGHT TO MEDICAL ADVICE AND CARE.
6	(a) IN GENERAL.—Part 7 of subtitle B of title I of
7	the Employee Retirement Income Security Act of 1974 (29
8	U.S.C. 1181 et seq.) is amended—
9	(1) by redesignating subpart C as subpart D ;
10	and
11	(2) by inserting after subpart B the following:

1	"Subpart C—Patient Right to Medical Advice and
2	Care
3	"SEC. 721. PATIENT ACCESS TO EMERGENCY MEDICAL
4	CARE.
5	"(a) Coverage of Emergency Care.—
б	"(1) IN GENERAL.—To the extent that the group
7	health plan (other than a fully insured group health
8	plan) provides coverage for benefits consisting of
9	emergency medical care (as defined in subsection (c))
10	or emergency ambulance services, except for items or
11	services specifically excluded—
12	"(A) the plan shall provide coverage for
13	benefits, without requiring preauthorization, for
14	emergency medical screening examinations or
15	emergency ambulance services, to the extent that
16	a prudent layperson, who possesses an average
17	knowledge of health and medicine, would deter-
18	mine such examinations or emergency ambu-
19	lance services to be necessary to determine wheth-
20	er emergency medical care (as so defined) is nec-
21	essary; and
22	"(B) the plan shall provide coverage for
23	benefits, without requiring preauthorization, for
24	additional emergency medical care to stabilize
25	an emergency medical condition following an
26	emergency medical screening examination (if de-

1	termined necessary under subparagraph (A)),
2	pursuant to the definition of stabilize under sec-
3	tion 1867(e)(3) of the Social Security Act (42
4	$U.S.C. \ 1395 dd(e)(3)).$
5	"(2) Reimbursement for care to maintain
6	MEDICAL STABILITY.—
7	"(A) IN GENERAL.—In the case of services
8	provided to a participant or beneficiary by a
9	nonparticipating provider in order to maintain
10	the medical stability of the participant or bene-
11	ficiary, the group health plan involved shall pro-
12	vide for reimbursement with respect to such serv-
13	ices if—
14	"(i) coverage for services of the type
15	furnished is available under the group
16	health plan;
17	"(ii) the services were provided for care
18	related to an emergency medical condition
19	and in an emergency department in order
20	to maintain the medical stability of the
21	participant or beneficiary; and
22	"(iii) the nonparticipating provider
23	contacted the plan regarding approval for
24	such services.

1	"(B) FAILURE TO RESPOND.—If a group
2	health plan fails to respond within 1 hours of
3	being contacted in accordance with subpara-
4	graph (A)(iii), then the plan shall be liable for
5	the cost of services provided by the nonpartici-
6	pating provider in order to maintain the sta-
7	bility of the participant or beneficiary.
8	"(C) LIMITATION.—The liability of a group
9	health plan to provide reimbursement under sub-
10	paragraph (A) shall terminate when the plan
11	has contacted the nonparticipating provider to
12	arrange for discharge or transfer.
13	"(D) LIABILITY OF PARTICIPANT.—A par-
14	ticipant or beneficiary shall not be liable for the
15	costs of services to which subparagraph (A) in an
16	amount that exceeds the amount of liability that
17	would be incurred if the services were provided
18	by a participating health care provider with
19	prior authorization by the plan.
20	"(b) IN-NETWORK UNIFORM COSTS-SHARING AND
21	Out-of-Network Care.—
22	"(1) IN-NETWORK UNIFORM COST-SHARING.—
23	Nothing in this section shall be construed as pre-
24	venting a group health plan (other than a fully in-
25	sured group health plan) from imposing any form of

1	cost-sharing applicable to any participant or bene-
2	ficiary (including coinsurance, copayments,
3	deductibles, and any other charges) in relation to cov-
4	erage for benefits described in subsection (a), if such
5	form of cost-sharing is uniformly applied under such
6	plan, with respect to similarly situated participants
7	and beneficiaries, to all benefits consisting of emer-
8	gency medical care (as defined in subsection (c)) pro-
9	vided to such similarly situated participants and
10	beneficiaries under the plan, and such cost-sharing is
11	disclosed in accordance with section 714.
12	"(2) Out-of-network care.—If a group health
13	plan (other than a fully insured group health plan)
14	provides any benefits with respect to emergency med-
15	ical care (as defined in subsection (c)), the plan shall
16	cover emergency medical care under the plan in a
17	manner so that, if such care is provided to a partici-
18	pant or beneficiary by a nonparticipating health care
19	provider, the participant or beneficiary is not liable
20	for amounts that exceed any form of cost-sharing (in-
21	cluding co-insurance, co-payments, deductibles, and
22	any other charges) that would be incurred if the serv-
23	ices were provided by a participating provider.
24	"(c) Definition of Emergency Medical Care.—In
25	this section:

1	"(1) IN GENERAL.—The term 'emergency medical
2	care' means, with respect to a participant or bene-
3	ficiary under a group health plan (other than a fully
4	insured group health plan), covered inpatient and
5	outpatient services that—
6	"(A) are furnished by any provider, includ-
7	ing a nonparticipating provider, that is quali-
8	fied to furnish such services; and
9	``(B) are needed to evaluate or stabilize (as
10	such term is defined in section $1867(e)(3)$ of the
11	Social Security Act (42 U.S.C. $1395dd$)(e)(3))
12	an emergency medical condition (as defined in
13	paragraph (2)).
14	"(2) Emergency medical condition.—The
15	term 'emergency medical condition' means a medical
16	condition manifesting itself by acute symptoms of suf-
17	ficient severity (including severe pain) such that a
18	prudent layperson, who possesses an average knowl-
19	edge of health and medicine, could reasonably expect
20	the absence of immediate medical attention to result
21	in—
22	"(A) placing the health of the participant or
23	beneficiary (or, with respect to a pregnant
24	woman, the health of the woman or her unborn
25	child) in serious jeopardy,

1	"(B) serious impairment to bodily func-
2	tions, or
-3	"(C) serious dysfunction of any bodily
4	organ or part.
5	"SEC. 722. OFFERING OF CHOICE OF COVERAGE OPTIONS.
6	"(a) Requirement.—
7	"(1) Offering of point-of-service coverage
8	OPTION.—Except as provided in paragraph (2), if a
9	group health plan (other than a fully insured group
10	health plan) provides coverage for benefits only
11	through a defined set of participating health care pro-
12	fessionals, the plan shall offer the participant the op-
13	tion to purchase point-of-service coverage (as defined
14	in subsection (b)) for all such benefits for which cov-
15	erage is otherwise so limited. Such option shall be
16	made available to the participant at the time of en-
17	rollment under the plan and at such other times as
18	the plan offers the participant a choice of coverage op-
19	tions.
20	"(2) Exception in case of lack of avail-

20 (2) EXCEPTION IN CASE OF LACK OF AVAIL21 ABILITY.—Paragraph (1) shall not apply with respect
22 to a group health plan (other than a fully insured
23 group health plan) if care relating to the point-of24 service coverage would not be available and accessible
25 to the participant with reasonable promptness (con-

sistent with section 1301(b)(4) of the Public Health
 Service Act (42 U.S.C. 300e(b)(4))).

3 "(b) POINT-OF-SERVICE COVERAGE DEFINED.—In
4 this section, the term 'point-of-service coverage' means, with
5 respect to benefits covered under a group health plan (other
6 than a fully insured group health plan), coverage of such
7 benefits when provided by a nonparticipating health care
8 professional.

9 "(c) Small Employer Exemption.—

"(1) IN GENERAL.—This section shall not apply
to any group health plan (other than a fully insured
group health plan) of a small employer.

13 "(2) SMALL EMPLOYER.—For purposes of para-14 graph (1), the term 'small employer' means, in con-15 nection with a group health plan (other than a fully 16 insured group health plan) with respect to a calendar 17 year and a plan year, an employer who employed an 18 average of at least 2 but not more than 50 employees 19 on business days during the preceding calendar year 20 and who employs at least 2 employees on the first day 21 of the plan year. For purposes of this paragraph, the 22 provisions of subparagraph (C) of section 712(c)(1)23 shall apply in determining employer size.

24 "(d) RULE OF CONSTRUCTION.—Nothing in this sec25 tion shall be construed—

1	"(1) as requiring coverage for benefits for a par-
2	ticular type of health care professional;
3	"(2) as requiring an employer to pay any costs
4	as a result of this section or to make equal contribu-
5	tions with respect to different health coverage options;
6	"(3) as preventing a group health plan (other
7	than a fully insured group health plan) from impos-
8	ing higher premiums or cost-sharing on a participant
9	for the exercise of a point-of-service coverage option;
10	or
11	"(4) to require that a group health plan (other
12	than a fully insured group health plan) include cov-
13	erage of health care professionals that the plan ex-
14	cludes because of fraud, quality of care, or other simi-
15	lar reasons with respect to such professionals.
16	"SEC. 723. PATIENT ACCESS TO OBSTETRIC AND GYNECO-
17	LOGICAL CARE.
18	"(a) General Rights.—
19	"(1) WAIVER OF PLAN REFERRAL REQUIRE-
20	MENT.—If a group health plan described in subsection
21	(b) requires a referral to obtain coverage for specialty
22	care, the plan shall waive the referral requirement in
23	the case of a female participant or beneficiary who
24	seeks coverage for obstetrical care and related follow-

1	up obstetrical care or routine gynecological care (such
2	as preventive gynecological care).
3	"(2) Related routine care.—With respect to
4	a participant or beneficiary described in paragraph
5	(1), a group health plan described in subsection (b)
6	shall treat the ordering of other routine care that is
7	related to routine gynecologic care, by a physician
8	who specializes in obstetrics and gynecology as the
9	authorization of the primary care provider for such
10	other care.
11	"(b) Application of Section.—A group health plan
12	described in this subsection is a group health plan (other
13	than a fully insured group health plan), that—
14	"(1) provides coverage for obstetric care (such as
15	pregnancy-related services) or routine gynecologic
16	care (such as preventive women's health examina-
17	tions); and
18	"(2) requires the designation by a participant or
19	beneficiary of a participating primary care provider
20	who is not a physician who specializes in obstetrics
21	or gynecology.
22	"(c) Rules of Construction.—Nothing in this sec-
23	tion shall be construed—
24	"(1) as waiving any coverage requirement relat-
25	ing to medical necessity or appropriateness with re-

1	spect to the coverage of obstetric or gynecologic care
2	described in subsection (a);
3	"(2) to preclude the plan from requiring that the
4	physician who specializes in obstetrics or gynecology
5	notify the designated primary care provider or the
6	plan of treatment decisions;
7	"(3) to preclude a group health plan from allow-
8	ing health care professionals other than physicians to
9	provide routine obstetric or routine gynecologic care;
10	OT
11	"(4) to preclude a group health plan from per-
12	mitting a physician who specializes in obstetrics and
13	gynecology from being a primary care provider under
14	the plan.
15	"SEC. 724. PATIENT ACCESS TO PEDIATRIC CARE.
16	"(a) IN GENERAL.—In the case of a group health plan
17	(other than a fully insured group health plan) that provides
18	coverage for routine pediatric care and that requires the
19	designation by a participant or beneficiary of a partici-
20	pating primary care provider, if the designated primary
21	care provider is not a physician who specializes in
22	pediatrics—
22	(1) the effect of an end of the second se

23 "(1) the plan may not require authorization or
24 referral by the primary care provider in order for a

1	participant or beneficiary to obtain coverage for rou-
2	tine pediatric care; and
3	"(2) the plan shall treat the ordering of other
4	routine care related to routine pediatric care by such
5	a specialist as having been authorized by the des-
6	ignated primary care provider.
7	"(b) Rules of Construction.—Nothing in sub-
8	section (a) shall be construed—
9	"(1) as waiving any coverage requirement relat-
10	ing to medical necessity or appropriateness with re-
11	spect to the coverage of any pediatric care provided
12	to, or ordered for, a participant or beneficiary;
13	"(2) to preclude a group health plan from re-
14	quiring that a specialist described in subsection (a)
15	notify the designated primary care provider or the
16	plan of treatment decisions; or
17	"(3) to preclude a group health plan from allow-
18	ing health care professionals other than physicians to
19	provide routine pediatric care.
20	"SEC. 725. TIMELY ACCESS TO SPECIALISTS.
21	"(a) TIMELY ACCESS.—
22	"(1) IN GENERAL.—A group health plan (other
23	than a fully insured group health plan) shall ensure
24	that participants and beneficiaries have timely, in
25	accordance with the medical exigencies of the case, ac-

1	cess to primary and specialty health care profes-
2	sionals who are appropriate to the condition of the
3	participant or beneficiary, when such care is covered
4	under the plan. Such access may be provided through
5	contractual arrangements with specialized providers
6	outside of the network of the plan.
7	"(2) RULE OF CONSTRUCTION.—Nothing in
8	paragraph (1) shall be construed—
9	"(A) to require the coverage under a group
10	health plan of particular benefits or services or
11	to prohibit a plan from including providers only
12	to the extent necessary to meet the needs of the
13	plan's participants or beneficiaries or from es-
14	tablishing any measure designed to maintain
15	quality and control costs consistent with the re-
16	sponsibilities of the plan; or
17	"(B) to override any State licensure or
18	scope-of-practice law.
19	"(b) TREATMENT PLANS.—
20	"(1) IN GENERAL.—Nothing in this section shall
21	be construed to prohibit a group health plan (other
22	than a fully insured group health plan) from requir-
23	ing that specialty care be provided pursuant to a
24	treatment plan so long as the treatment plan is—

1	((A) developed by the specialist, in con-
2	sultation with the case manager or primary care
3	provider, and the participant or beneficiary;
4	"(B) approved by the plan in a timely
5	manner in accordance with the medical exigen-
6	cies of the case; and
7	"(C) in accordance with the applicable (C)
8	quality assurance and utilization review stand-
9	ards of the plan.
10	"(2) NOTIFICATION.—Nothing in paragraph (1)
11	shall be construed as prohibiting a plan from requir-
12	ing the specialist to provide the case manager or pri-
13	mary care provider with regular updates on the spe-
14	cialty care provided, as well as all other necessary
15	medical information.
16	"(c) Referrals.—Nothing in this section shall be
17	construed to prohibit a plan from requiring an authoriza-
18	tion by the case manager or primary care provider of the
19	participant or beneficiary in order to obtain coverage for

20 specialty services so long as such authorization is for an
21 adequate number of referrals.

"(d) SPECIALTY CARE DEFINED.—For purposes of this
subsection, the term 'specialty care' means, with respect to
a condition, care and treatment provided by a health care
practitioner, facility, or center (such as a center of excel-

lence) that has adequate expertise (including age-appro priate expertise) through appropriate training and experi ence.

4 "SEC. 726. CONTINUITY OF CARE.

5 "(a) IN GENERAL.—

"(1) TERMINATION OF PROVIDER.—If a contract 6 7 between a group health plan (other than a fully in-8 sured group health plan) and a health care provider 9 is terminated (as defined in paragraph (2)), or bene-10 fits or coverage provided by a health care provider are 11 terminated because of a change in the terms of pro-12 vider participation in such group health plan, and an individual who is a participant or beneficiary in 13 14 the plan is undergoing a course of treatment from the 15 provider at the time of such termination, the plan shall-16

17 "(A) notify the individual on a timely basis
18 of such termination;

19 "(B) provide the individual with an oppor20 tunity to notify the plan of a need for transi21 tional care; and

"(C) in the case of termination described in
paragraph (2), (3), or (4) of subsection (b), and
subject to subsection (c), permit the individual to
continue or be covered with respect to the course

1	of treatment with the provider's consent during
2	a transitional period (as provided under sub-
3	section (b)).
4	"(2) TERMINATED.—In this section, the term
5	'terminated' includes, with respect to a contract, the
6	expiration or nonrenewal of the contract by the group
7	health plan, but does not include a termination of the
8	contract by the plan for failure to meet applicable
9	quality standards or for fraud.
10	"(3) CONTRACTS.—For purposes of this section,
11	the term 'contract between a group health plan (other
12	than a fully insured group health plan) and a health
13	care provider' shall include a contract between such
14	a plan and an organized network of providers.
15	"(b) Transitional Period.—
16	"(1) GENERAL RULE.—Except as provided in
17	paragraph (3), the transitional period under this sub-
18	section shall permit the participant or beneficiary to
19	extend the coverage involved for up to 90 days from
20	the date of the notice described in subsection $(a)(1)(A)$
21	of the provider's termination.
22	"(2) INSTITUTIONAL CARE.—Subject to para-
23	graph (1), the transitional period under this sub-
24	section for institutional or inpatient care from a pro-
25	vider shall extend until the discharge or termination

1	of the period of institutionalization and also shall in-
2	clude institutional care provided within a reasonable
3	time of the date of termination of the provider status
4	if the care was scheduled before the date of the an-
5	nouncement of the termination of the provider status
6	under subsection $(a)(1)(A)$ or if the individual on
7	such date was on an established waiting list or other-
8	wise scheduled to have such care.
9	"(3) Pregnancy.—Notwithstanding paragraph
10	(1), if—
11	``(A) a participant or beneficiary has en-
12	tered the second trimester of pregnancy at the
13	time of a provider's termination of participa-
14	tion; and
15	``(B) the provider was treating the preg-
16	nancy before the date of the termination;
17	the transitional period under this subsection with re-
18	spect to provider's treatment of the pregnancy shall
19	extend through the provision of post-partum care di-
20	rectly related to the delivery.
21	"(4) TERMINAL ILLNESS.—Notwithstanding
22	paragraph (1), if—
23	"(A) a participant or beneficiary was deter-
24	mined to be terminally ill (as determined under
25	section $1861(dd)(3)(A)$ of the Social Security

1	Act) prior to a provider's termination of partici-
2	pation; and
3	"(B) the provider was treating the terminal
4	illness before the date of termination;
5	the transitional period under this subsection shall be
6	for care directly related to the treatment of the ter-
7	minal illness and shall extend for the remainder of
8	the individual's life for such care.
9	"(c) Permissible Terms and Conditions.—A group
10	health plan (other than a fully insured group health plan)
11	may condition coverage of continued treatment by a pro-
12	vider under subsection $(a)(1)(C)$ upon the provider agreeing
13	to the following terms and conditions:
14	"(1) The provider agrees to accept reimburse-
15	ment from the plan and individual involved (with re-
16	spect to cost-sharing) at the rates applicable prior to
17	the start of the transitional period as payment in full
18	(or at the rates applicable under the replacement plan
19	after the date of the termination of the contract with
20	the group health plan) and not to impose cost-sharing
21	with respect to the individual in an amount that
22	would exceed the cost-sharing that could have been
23	imposed if the contract referred to in subsection $(a)(1)$
24	had not been terminated.

1	"(2) The provider agrees to adhere to the quality
2	assurance standards of the plan responsible for pay-
3	ment under paragraph (1) and to provide to such
4	plan necessary medical information related to the
5	care provided.
6	"(3) The provider agrees otherwise to adhere to
7	such plan's policies and procedures, including proce-
8	dures regarding referrals and obtaining prior author-
9	ization and providing services pursuant to a treat-
10	ment plan (if any) approved by the plan.
11	"(d) Rule of Construction.—Nothing in this sec-
12	tion shall be construed to require the coverage of benefits
13	which would not have been covered if the provider involved
14	remained a participating provider.
15	"(e) DEFINITION.—In this section, the term health
16	care provider' or 'provider' means—
17	"(1) any individual who is engaged in the deliv-
18	ery of health care services in a State and who is re-
19	quired by State law or regulation to be licensed or
20	certified by the State to engage in the delivery of such
21	services in the State; and
22	"(2) any entity that is engaged in the delivery
23	
	of health care services in a State and that, if it is re-

certified by the State to engage in the delivery of such
 services in the State, is so licensed.

3 "(f) COMPREHENSIVE STUDY OF COST, QUALITY AND
4 COORDINATION OF COVERAGE FOR PATIENTS AT THE END
5 OF LIFE.—

6 "(1) Study by the medicare payment advi-SORY COMMISSION.—The Medicare Payment Advisory 7 Commission shall conduct a study of the costs and 8 9 patterns of care for persons with serious and complex 10 conditions and the possibilities of improving upon 11 that care to the degree it is triggered by the current 12 category of terminally ill as such term is used for 13 purposes of section 1861(dd) of the Social Security 14 Act (relating to hospice benefits) or of utilizing care 15 in other payment settings in Medicare.

"(2) AGENCY FOR HEALTH CARE POLICY AND RESEARCH.—The Agency for Health Care Policy and
Research shall conduct studies of the possible thresholds for major conditions causing serious and complex
illness, their administrative parameters and feasibility, and their impact upon costs and quality.

22 "(3) HEALTH CARE FINANCING ADMINISTRA23 TION.—The Health Care Financing Administration
24 shall conduct studies of the merits of applying similar
25 thresholds in Medicare+Choice programs, including

adapting risk adjustment methods to account for this
 category.

"(4) Initial report.—

3

4 "(A) IN GENERAL.—Not later than 12 months after the date of enactment of this sec-5 6 tion, the Medicare Payment Advisory Commis-7 sion and the Agency for Health Care Policy and 8 Research shall each prepare and submit to the 9 Committee on Health, Education, Labor and Pensions of the Senate a report concerning the 10 11 results of the studies conducted under para-12 graphs (1) and (2), respectively.

"(B) COPY TO SECRETARY.—Concurrent
with the submission of the reports under subparagraph (A), the Medicare Payment Advisory
Commission and the Agency for health Care Policy and Research shall transmit a copy of the reports under such subparagraph to the Secretary.
"(5) FINAL REPORT.—

20 "(A) CONTRACT WITH INSTITUTE OF MEDI21 CINE.—Not later than 1 year after the submis22 sion of the reports under paragraph (4), the Sec23 retary of Health and Human Services shall con24 tract with the Institute of Medicine to conduct a
25 study of the practices and their effects arising

3 "(B) REPORT.—Not later than 1 year after
4 the date of the execution of the contract referred
5 to in subparagraph (A), the Institute of Medicine
6 shall prepare and submit to the Committee on
7 Health, Education, Labor and Pensions of the
8 Senate a report concerning the study conducted
9 pursuant to such contract.

10 "(6) FUNDING.—From funds appropriated to the
11 Department of Health and Human Services, the Sec12 retary of Health and Human Services shall make
13 available such funds as the Secretary determines is
14 necessary to carry out this subsection.

15 "SEC. 727. PROTECTION OF PATIENT-PROVIDER COMMU 16 NICATIONS.

17 "(a) IN GENERAL.—Subject to subsection (b), a group 18 health plan (other than a fully insured group health plan 19 and in relation to a participant or beneficiary) shall not prohibit or otherwise restrict a health care professional from 20 21 advising such a participant or beneficiary who is a patient 22 of the professional about the health status of the participant 23 or beneficiary or medical care or treatment for the condition 24 or disease of the participant or beneficiary, regardless of whether coverage for such care or treatment are provided 25

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under the contract, if the professional is acting within the
 lawful scope of practice.

3 "(b) RULE OF CONSTRUCTION.—Nothing in this sec4 tion shall be construed as requiring a group health plan
5 (other than a fully insured group health plan) to provide
6 specific benefits under the terms of such plan.

7 "SEC. 728. PATIENT'S RIGHT TO PRESCRIPTION DRUGS.

8 "To the extent that a group health plan (other than 9 a fully insured group health plan) provides coverage for 10 benefits with respect to prescription drugs, and limits such 11 coverage to drugs included in a formulary, the plan shall— 12 "(1) ensure the participation of physicians and 13 pharmacists in developing and reviewing such for-14 mulary; and

"(2) in accordance with the applicable quality
assurance and utilization review standards of the
plan, provide for exceptions from the formulary limitation when a non-formulary alternative is medically
necessary and appropriate.

20 "SEC. 729. SELF-PAYMENT FOR BEHAVIORAL HEALTH CARE
21 SERVICES.

22 "(a) IN GENERAL.—A group health plan (other than
23 a fully insured group health plan) may not—

24 "(1) prohibit or otherwise discourage a partici25 pant or beneficiary from self-paying for behavioral

1	health care services once the plan has denied coverage
2	for such services; or
3	"(2) terminate a health care provider because
4	such provider permits participants or beneficiaries to
5	self-pay for behavioral health care services—
6	"(A) that are not otherwise covered under
7	the plan; or
8	"(B) for which the group health plan pro-
9	vides limited coverage, to the extent that the
10	group health plan denies coverage of the services.
11	"(b) RULE OF CONSTRUCTION.—Nothing in subsection
12	(a)(2)(B) shall be construed as prohibiting a group health
13	plan from terminating a contract with a health care pro-
14	vider for failure to meet applicable quality standards or
15	for fraud.
16	"SEC. 730. COVERAGE FOR INDIVIDUALS PARTICIPATING IN
17	APPROVED CANCER CLINICAL TRIALS.
18	"(a) COVERAGE.—
19	"(1) IN GENERAL.—If a group health plan (other
20	than a fully insured group health plan) provides cov-
21	erage to a qualified individual (as defined in sub-
22	section (b)), the plan—
23	"(A) may not deny the individual partici-
24	pation in the clinical trial referred to in sub-
25	section $(b)(2);$

"(B) subject to subsections (b), (c), and (d)
may not deny (or limit or impose additional
conditions on) the coverage of routine patient
costs for items and services furnished in connec-
tion with participation in the trial; and
``(C) may not discriminate against the in-
dividual on the basis of the participant's or
beneficiaries participation in such trial.
"(2) Exclusion of certain costs.—For pur-
poses of paragraph $(1)(B)$, routine patient costs do
not include the cost of the tests or measurements con-
ducted primarily for the purpose of the clinical trial
involved.
"(3) Use of in-network providers.—If one or
more participating providers is participating in a
clinical trial, nothing in paragraph (1) shall be con-
strued as preventing a plan from requiring that a
qualified individual participate in the trial through
such a participating provider if the provider will ac-
cept the individual as a participant in the trial.
cept the individual as a participant in the trial. "(b) QUALIFIED INDIVIDUAL DEFINED.—For purposes
"(b) Qualified Individual Defined.—For purposes

1	((1)(A) The individual has been diagnosed with
2	cancer for which no standard treatment is effective.
3	``(B) The individual is eligible to participate in
4	an approved clinical trial according to the trial pro-
5	tocol with respect to treatment of such illness.
6	``(C) The individual's participation in the trial
7	offers meaningful potential for significant clinical
8	benefit for the individual.
9	"(2) Either—
10	"(A) the referring physician is a partici-
11	pating health care professional and has con-
12	cluded that the individual's participation in
13	such trial would be appropriate based upon the
14	individual meeting the conditions described in
15	paragraph (1); or
16	"(B) the participant or beneficiary provides
17	medical and scientific information establishing
18	that the individual's participation in such trial
19	would be appropriate based upon the individual
20	meeting the conditions described in paragraph
21	(1).
22	"(c) PAYMENT.—
23	"(1) IN GENERAL.—Under this section a group
24	health plan (other than a fully insured group health
25	plan) shall provide for payment for routine patient

1	costs described in subsection $(a)(2)$ but is not required
2	to pay for costs of items and services that are reason-
3	ably expected to be paid for by the sponsors of an ap-
4	proved clinical trial.
5	"(2) Standards for determining routine
6	PATIENT COSTS ASSOCIATED WITH CLINICAL TRIAL
7	PARTICIPATION.—
8	"(A) IN GENERAL.—The Secretary shall es-
9	tablish, on an expedited basis and using a nego-
10	tiated rulemaking process under subchapter III
11	of chapter 5 of title 5, United States Code, stand-
12	ards relating to the coverage of routine patient
13	costs for individuals participating in clinical
14	trials that group health plans must meet under
15	this section.
16	"(B) FACTORS.—In establishing routine pa-
17	tient cost standards under subparagraph (A), the
18	Secretary shall consult with interested parties
19	and take into account —
20	"(i) quality of patient care;
21	"(ii) routine patient care costs versus
22	costs associated with the conduct of clinical
23	trials, including unanticipated patient care
24	costs as a result of participation in clinical
25	trials; and

1	"(iii) previous and on-going studies re-
2	lating to patient care costs associated with
3	participation in clinical trials.

4 "(C) PUBLICATION OF NOTICE.—In car-5 rying out the rulemaking process under this 6 paragraph, the Secretary, after consultation with 7 organizations representing cancer patients. 8 health care practitioners, medical researchers, 9 employers, group health plans, manufacturers of 10 drugs, biologics and medical devices, medical 11 economists, hospitals, and other interested par-12 ties, shall publish notice provided for under sec-13 tion 564(a) of title 5, United States Code, by not 14 later than 45 days after the date of the enact-15 ment of this section.

"(D) TARGET DATE FOR PUBLICATION OF
RULE.—As part of the notice under subparagraph (C), and for purposes of this paragraph,
the 'target date for publication' (referred to in
section 564(a)(5) of such title 5) shall be June
30, 2000.

22 "(E) ABBREVIATED PERIOD FOR SUBMIS23 SION OF COMMENTS.—In applying section 564(c)
24 of such title 5 under this paragraph, '15 days'
25 shall be substituted for '30 days'.

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1	"(F) Appointment of negotiated rule-
2	MAKING COMMITTEE AND FACILITATOR.—The
3	Secretary shall provide for—
4	"(i) the appointment of a negotiated
5	$rule making \ committee \ under \ section \ 565(a)$
6	of such title 5 by not later than 30 days
7	after the end of the comment period pro-
8	vided for under section $564(c)$ of such title
9	5 (as shortened under subparagraph (E)),
10	and
11	"(ii) the nomination of a facilitator
12	under section $566(c)$ of such title 5 by not
13	later than 10 days after the date of appoint-
14	ment of the committee.
15	"(G) Preliminary committee report.—
16	The negotiated rulemaking committee appointed
17	under subparagraph (F) shall report to the Sec-
18	retary, by not later than March 29, 2000, re-
19	garding the committee's progress on achieving a
20	consensus with regard to the rulemaking pro-
21	ceeding and whether such consensus is likely to
22	occur before 1 month before the target date for
23	publication of the rule. If the committee reports
24	that the committee has failed to make significant
25	progress towards such consensus or is unlikely to

1	reach such consensus by the target date, the Sec-
2	retary may terminate such process and provide
3	for the publication of a rule under this para-
4	graph through such other methods as the Sec-
5	retary may provide.
6	"(H) FINAL COMMITTEE REPORT.—If the
7	committee is not terminated under subparagraph
8	(G), the rulemaking committee shall submit a re-
9	port containing a proposed rule by not later
10	than 1 month before the target date of publica-
11	tion.
12	"(I) FINAL EFFECT.—The Secretary shall
13	publish a rule under this paragraph in the Fed-
14	eral Register by not later than the target date of
15	publication.
16	"(J) Publication of rule after public
17	COMMENT.—The Secretary shall provide for con-
18	sideration of such comments and republication of
19	such rule by not later than 1 year after the tar-
20	get date of publication.
21	"(K) EFFECTIVE DATE.—The provisions of
22	this paragraph shall apply to group health plans
23	(other than a fully insured group health plan)
24	for plan years beginning on or after January 1,
25	2001.

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"(3) PAYMENT RATE.—In the case of covered
items and services provided by—
"(A) a participating provider, the payment
rate shall be at the agreed upon rate, or
"(B) a nonparticipating provider, the pay-
ment rate shall be at the rate the plan would
normally pay for comparable services under sub-
paragraph (A).
"(d) Approved Clinical Trial Defined.—
"(1) IN GENERAL.—In this section, the term 'ap-
proved clinical trial' means a cancer clinical research
study or cancer clinical investigation approved and
funded (which may include funding through in-kind
contributions) by one or more of the following:
"(A) The National Institutes of Health.
"(B) A cooperative group or center of the
National Institutes of Health.
"(C) Either of the following if the condi-
tions described in paragraph (2) are met:
"(i) The Department of Veterans Af-
fairs.
"(ii) The Department of Defense.
"(2) Conditions for departments.—The con-
ditions described in this paragraph, for a study or in-
vestigation conducted by a Department, are that the

1	study or investigation has been reviewed and ap-
2	proved through a system of peer review that the Sec-
3	retary determines—
4	"(A) to be comparable to the system of peer
5	review of studies and investigations used by the
6	National Institutes of Health, and
7	``(B) assures unbiased review of the highest
8	scientific standards by qualified individuals who
9	have no interest in the outcome of the review.
10	"(e) CONSTRUCTION.—Nothing in this section shall be
11	construed to limit a plan's coverage with respect to clinical
12	trials.
13	"(f) Plan Satisfaction of Certain Requirements;
14	Responsibilities of Fiduciaries.—
15	"(1) IN GENERAL.—For purposes of this section,
16	insofar as a group health plan provides benefits in the
17	form of health insurance coverage through a health in-
18	surance issuer, the plan shall be treated as meeting
19	the requirements of this section with respect to such
20	benefits and not be considered as failing to meet such
21	requirements because of a failure of the issuer to meet
22	such requirements so long as the plan sponsor or its
23	representatives did not cause such failure by the
24	issuer.

1	"(2) CONSTRUCTION.—Nothing in this section
2	shall be construed to affect or modify the responsibil-
3	ities of the fiduciaries of a group health plan under
4	part 4 of subtitle B.
5	"(g) Study and Report.—
6	"(1) STUDY.—The Secretary shall study the im-
7	pact on group health plans for covering routine pa-
8	tient care costs for individuals who are entitled to
9	benefits under this section and who are enrolled in an
10	approved cancer clinical trial program.
11	"(2) Report to congress.—Not later than
12	January 1, 2005, the Secretary shall submit a report
13	to Congress that contains an assessment of—
14	"(A) any incremental cost to group health
15	plans resulting from the provisions of this sec-
15	plans resulting from the provisions of this sec-
15 16	plans resulting from the provisions of this sec- tion;
15 16 17	plans resulting from the provisions of this sec- tion; "(B) a projection of expenditures to such
15 16 17 18	plans resulting from the provisions of this sec- tion; "(B) a projection of expenditures to such plans resulting from this section; and
15 16 17 18 19	<pre>plans resulting from the provisions of this sec- tion;</pre>
15 16 17 18 19 20	<pre>plans resulting from the provisions of this sec- tion;</pre>
15 16 17 18 19 20 21	<pre>plans resulting from the provisions of this sec- tion;</pre>
 15 16 17 18 19 20 21 22 	plans resulting from the provisions of this sec- tion; (B) a projection of expenditures to such plans resulting from this section; and (C) any impact on premiums resulting from this section. *SEC. 730A. PROHIBITING DISCRIMINATION AGAINST PRO- VIDERS.
1 provider who is acting within the scope of the provider's license or certification under applicable State law, solely 2 on the basis of such license or certification. This subsection 3 4 shall not be construed as requiring the coverage under a 5 plan of particular benefits or services or to prohibit a plan from including providers only to the extent necessary to 6 7 meet the needs of the plan's participants and beneficiaries 8 or from establishing any measure designed to maintain 9 quality and control costs consistent with the responsibilities 10 of the plan.

11 "(b) NO REQUIREMENT FOR ANY WILLING PRO-12 VIDER.—Nothing in this section shall be construed as re-13 quiring a group health plan that offers network coverage 14 to include for participation every willing provider or health 15 professional who meets the terms and conditions of the plan.

16 "SEC. 730B. GENERALLY APPLICABLE PROVISION.

17 "In the case of a group health plan that provides bene18 fits under 2 or more coverage options, the requirements of
19 this subpart shall apply separately with respect to each cov20 erage option.".

21 (b) RULE WITH RESPECT TO CERTAIN PLANS.—

(1) IN GENERAL.—Notwithstanding any other
provision of law, health insurance issuers may offer,
and eligible individuals may purchase, high deductible health plans described in section 220(c)(2)(A) of

1	the Internal Revenue Code of 1986. Effective for the
2	4-year period beginning on the date of the enactment
3	of this Act, such health plans shall not be required to
4	provide payment for any health care items or services
5	that are exempt from the plan's deductible.
6	(2) Existing state laws.—A State law relat-
7	ing to payment for health care items and services in
8	effect on the date of enactment of this Act that is pre-
9	empted under paragraph (1), shall not apply to high
10	deductible health plans after the expiration of the 4-
11	year period described in such paragraph unless the
12	State reenacts such law after such period.
13	(c) DEFINITION.—Section 733(a) of the Employee Re-
15	
13	tirement Income Security Act of 1974 (42 U.S.C. 1191(a))
14	tirement Income Security Act of 1974 (42 U.S.C. 1191(a))
14 15	tirement Income Security Act of 1974 (42 U.S.C. 1191(a)) is amended by adding at the end the following:
14 15 16	tirement Income Security Act of 1974 (42 U.S.C. 1191(a)) is amended by adding at the end the following: "(3) FULLY INSURED GROUP HEALTH PLAN.—
14 15 16 17	tirement Income Security Act of 1974 (42 U.S.C. 1191(a)) is amended by adding at the end the following: "(3) FULLY INSURED GROUP HEALTH PLAN.— The term 'fully insured group health plan' means a
14 15 16 17 18	tirement Income Security Act of 1974 (42 U.S.C. 1191(a)) is amended by adding at the end the following: "(3) FULLY INSURED GROUP HEALTH PLAN.— The term 'fully insured group health plan' means a group health plan where benefits under the plan are
14 15 16 17 18 19	tirement Income Security Act of 1974 (42 U.S.C. 1191(a)) is amended by adding at the end the following: "(3) FULLY INSURED GROUP HEALTH PLAN.— The term 'fully insured group health plan' means a group health plan where benefits under the plan are provided pursuant to the terms of an arrangement be-
 14 15 16 17 18 19 20 	tirement Income Security Act of 1974 (42 U.S.C. 1191(a)) is amended by adding at the end the following: "(3) FULLY INSURED GROUP HEALTH PLAN.— The term 'fully insured group health plan' means a group health plan where benefits under the plan are provided pursuant to the terms of an arrangement be- tween a group health plan and a health insurance
 14 15 16 17 18 19 20 21 	tirement Income Security Act of 1974 (42 U.S.C. 1191(a)) is amended by adding at the end the following: "(3) FULLY INSURED GROUP HEALTH PLAN.— The term 'fully insured group health plan' means a group health plan where benefits under the plan are provided pursuant to the terms of an arrangement be- tween a group health plan and a health insurance issuer and are guaranteed by the health insurance

1	(1) in the item relating to subpart C, by striking
2	"Subpart C" and inserting "Subpart D"; and
3	(2) by adding at the end of the items relating to
4	subpart B of part 7 of subtitle B of title I of such Act
5	the following new items:
	"SUBPART C-PATIENT RIGHT TO MEDICAL ADVICE AND CARE
	 "Sec. 721. Patient access to emergency medical care. "Sec. 722. Offering of choice of coverage options. "Sec. 723. Patient access to obstetric and gynecological care. "Sec. 724. Patient access to pediatric care. "Sec. 725. Timely access to specialists. "Sec. 726. Continuity of care. "Sec. 727. Protection of patient-provider communications. "Sec. 728. Patient's right to prescription drugs. "Sec. 729. Self-payment for behavioral health care services. "Sec. 730. Coverage for individuals participating in approved cancer clinical trials. "Sec. 730B. Generally applicable provision.".
6	SEC. 102. CONFORMING AMENDMENT TO THE INTERNAL
7	REVENUE CODE OF 1986.
8	(a) IN GENERAL.—Chapter 100 of the Internal Rev-
9	enue Code of 1986 is amended—
10	(1) by redesignating subchapter C as subchapter
11	D: and

12 (2) by inserting after subchapter B the following:

13 "Subchapter C—Patient Right to Medical

- 14 Advice and Care
 - "Sec. 9821. Patient access to emergency medical care.
 - "Sec. 9822. Offering of choice of coverage options.
 - "Sec. 9823. Patient access to obstetric and gynecological care.
 - "Sec. 9824. Patient access to pediatric care.
 - "Sec. 9825. Timely access to specialists.
 - "Sec. 9826. Continuity of care.
 - "Sec. 9827. Protection of patient-provider communications.
 - "Sec. 9828. Patient's right to prescription drugs.

"Sec. 9829. Self-payment for behavioral health care services.
"Sec. 9830. Coverage for individuals participating in approved cancer clinical trials.
"Sec. 9830A. Prohibiting discrimination against providers.

"Sec. 9830B. Generally applicable provision.

1 "SEC. 9821. PATIENT ACCESS TO EMERGENCY MEDICAL

CARE.

2

3 *"(a) COVERAGE OF EMERGENCY CARE.*—

4 "(1) IN GENERAL.—To the extent that the group
5 health plan (other than a fully insured group health
6 plan) provides coverage for benefits consisting of
7 emergency medical care (as defined in subsection (c))
8 or emergency ambulance services, except for items or
9 services specifically excluded—

"(A) the plan shall provide coverage for 10 11 benefits, without requiring preauthorization, for 12 emergency medical screening examinations or 13 emergency ambulance services, to the extent that 14 a prudent layperson, who possesses an average 15 knowledge of health and medicine, would deter-16 mine such examinations or emergency ambu-17 lance services to be necessary to determine wheth-18 er emergency medical care (as so defined) is nec-19 essary; and

20 "(B) the plan shall provide coverage for
21 benefits, without requiring preauthorization, for
22 additional emergency medical care to stabilize
23 an emergency medical condition following an

1	emergency medical screening examination (if de-
2	termined necessary under subparagraph (A)),
3	pursuant to the definition of stabilize under sec-
4	tion 1867(e)(3) of the Social Security Act (42
5	$U.S.C. \ 1395dd(e)(3)).$
6	"(2) Reimbursement for care to maintain
7	MEDICAL STABILITY.—
8	"(A) IN GENERAL.—In the case of services
9	provided to a participant or beneficiary by a
10	nonparticipating provider in order to maintain
11	the medical stability of the participant or bene-
12	ficiary, the group health plan involved shall pro-
13	vide for reimbursement with respect to such serv-
14	ices if—
15	((i) coverage for services of the type
16	furnished is available under the group
17	health plan;
18	"(ii) the services were provided for care
19	related to an emergency medical condition
20	and in an emergency department in order
21	to maintain the medical stability of the
22	participant or beneficiary; and
23	"(iii) the nonparticipating provider
24	contacted the plan regarding approval for
25	such services.

1	"(B) FAILURE TO RESPOND.—If a group
2	health plan fails to respond within 1 hours of
3	being contacted in accordance with subpara-
4	graph $(A)(iii)$, then the plan shall be liable for
5	the cost of services provided by the nonpartici-
6	pating provider in order to maintain the sta-
7	bility of the participant or beneficiary.
8	"(C) LIMITATION.—The liability of a group
9	health plan to provide reimbursement under sub-
10	paragraph (A) shall terminate when the plan
11	has contacted the nonparticipating provider to
12	arrange for discharge or transfer.
13	"(D) LIABILITY OF PARTICIPANT.—A par-
14	ticipant or beneficiary shall not be liable for the
15	costs of services to which subparagraph (A) in an
16	amount that exceeds the amount of liability that
17	would be incurred if the services were provided
18	by a participating health care provider with
19	prior authorization by the plan.
20	"(b) IN-NETWORK UNIFORM COSTS-SHARING AND
21	Out-of-Network Care.—
22	"(1) IN-NETWORK UNIFORM COST-SHARING.—
23	Nothing in this section shall be construed as pre-
24	venting a group health plan (other than a fully in-
25	sured group health plan) from imposing any form of

1	cost-sharing applicable to any participant or bene-
2	ficiary (including coinsurance, copayments,
3	deductibles, and any other charges) in relation to cov-
4	erage for benefits described in subsection (a), if such
5	form of cost-sharing is uniformly applied under such
6	plan, with respect to similarly situated participants
7	and beneficiaries, to all benefits consisting of emer-
8	gency medical care (as defined in subsection (c)) pro-
9	vided to such similarly situated participants and
10	beneficiaries under the plan, and such cost-sharing is
11	disclosed in accordance with section 9814.
12	"(2) Out-of-network care.—If a group health
13	plan (other than a fully insured group health plan)
14	provides any benefits with respect to emergency med-
15	ical care (as defined in subsection (c)), the plan shall
16	cover emergency medical care under the plan in a
17	manner so that, if such care is provided to a partici-
18	pant or beneficiary by a nonparticipating health care
19	provider, the participant or beneficiary is not liable
20	for amounts that exceed any form of cost-sharing (in-
21	cluding coinsurance, copayments, deductibles, and
22	any other charges) that would be incurred if the serv-
23	ices were provided by a participating provider.
24	"(c) Definition of Emergency Medical Care.—In
25	this section:

1	"(1) IN GENERAL.—The term 'emergency medical
2	care' means, with respect to a participant or bene-
3	ficiary under a group health plan (other than a fully
4	insured group health plan), covered inpatient and
5	outpatient services that—
6	"(A) are furnished by any provider, includ-
7	ing a nonparticipating provider, that is quali-
8	fied to furnish such services; and
9	``(B) are needed to evaluate or stabilize (as
10	such term is defined in section $1867(e)(3)$ of the
11	Social Security Act (42 U.S.C. $1395dd$)(e)(3))
12	an emergency medical condition (as defined in
13	paragraph (2)).
14	"(2) Emergency medical condition.—The
15	term 'emergency medical condition' means a medical
16	condition manifesting itself by acute symptoms of suf-
17	ficient severity (including severe pain) such that a
18	prudent layperson, who possesses an average knowl-
19	edge of health and medicine, could reasonably expect
20	the absence of immediate medical attention to result
21	in—
22	"(A) placing the health of the participant or
23	beneficiary (or, with respect to a pregnant
24	woman, the health of the woman or her unborn
25	child) in serious jeopardy,

1	``(B) serious impairment to bodily func-
2	tions, or
3	"(C) serious dysfunction of any bodily
4	organ or part.
5	"SEC. 9822. OFFERING OF CHOICE OF COVERAGE OPTIONS.
6	"(a) Requirement.—
7	"(1) Offering of point-of-service coverage
8	OPTION.—Except as provided in paragraph (2), if a
9	group health plan (other than a fully insured group
10	health plan) provides coverage for benefits only
11	through a defined set of participating health care pro-
12	fessionals, the plan shall offer the participant the op-
13	tion to purchase point-of-service coverage (as defined
14	in subsection (b)) for all such benefits for which cov-
15	erage is otherwise so limited. Such option shall be
16	made available to the participant at the time of en-
17	rollment under the plan and at such other times as
18	the plan offers the participant a choice of coverage op-
19	tions.
20	"(2) Exception in case of lack of avail-
21	ABILITY.—Paragraph (1) shall not apply with respect

22 to a group health plan (other than a fully insured 23 group health plan) if care relating to the point-of-24 service coverage would not be available and accessible 25 to the participant with reasonable promptness (consistent with section 1301(b)(4) of the Public Health
 Service Act (42 U.S.C. 300e(b)(4))).

3 "(b) POINT-OF-SERVICE COVERAGE DEFINED.—In
4 this section, the term 'point-of-service coverage' means, with
5 respect to benefits covered under a group health plan (other
6 than a fully insured group health plan), coverage of such
7 benefits when provided by a nonparticipating health care
8 professional.

9 "(c) SMALL EMPLOYER EXEMPTION.—

"(1) IN GENERAL.—This section shall not apply
to any group health plan (other than a fully insured
group health plan) of a small employer.

13 "(2) SMALL EMPLOYER.—For purposes of para-14 graph (1), the term 'small employer' means, in con-15 nection with a group health plan (other than a fully 16 insured group health plan) with respect to a calendar 17 year and a plan year, an employer who employed an 18 average of at least 2 but not more than 50 employees 19 on business days during the preceding calendar year 20 and who employs at least 2 employees on the first day 21 of the plan year. For purposes of this paragraph, the 22 provisions ofsubparagraph (C)ofsection 23 4980D(d)(2) shall apply in determining employer 24 size.

1	"(d) RULE OF CONSTRUCTION.—Nothing in this sec-
2	tion shall be construed—
3	"(1) as requiring coverage for benefits for a par-
4	ticular type of health care professional;
5	"(2) as requiring an employer to pay any costs
6	as a result of this section or to make equal contribu-
7	tions with respect to different health coverage options;
8	((3) as preventing a group health plan (other
9	than a fully insured group health plan) from impos-
10	ing higher premiums or cost-sharing on a participant
11	for the exercise of a point-of-service coverage option;
12	OT
13	"(4) to require that a group health plan (other
14	than a fully insured group health plan) include cov-
15	erage of health care professionals that the plan ex-
16	cludes because of fraud, quality of care, or other simi-
17	lar reasons with respect to such professionals.
18	"SEC. 9823. PATIENT ACCESS TO OBSTETRIC AND GYNECO-
19	LOGICAL CARE.
20	"(a) General Rights.—
21	"(1) WAIVER OF PLAN REFERRAL REQUIRE-
22	MENT.—If a group health plan described in subsection
23	(b) requires a referral to obtain coverage for specialty
24	care, the plan shall waive the referral requirement in
25	the case of a female participant or beneficiary who

1	seeks coverage for obstetrical care and related follow-
2	up obstetrical care or routine gynecological care (such
3	as preventive gynecological care).
4	"(2) Related routine care.—With respect to
5	a participant or beneficiary described in paragraph
6	(1), a group health plan described in subsection (b)
7	shall treat the ordering of other routine care that is
8	related to routine gynecologic care, by a physician
9	who specializes in obstetrics and gynecology as the
10	authorization of the primary care provider for such
11	other care.
12	"(b) Application of Section.—A group health plan
13	described in this subsection is a group health plan (other
13	described in this subsection is a group health plan (other
13 14	described in this subsection is a group health plan (other than a fully insured group health plan), that—
13 14 15	described in this subsection is a group health plan (other than a fully insured group health plan), that— "(1) provides coverage for obstetric care (such as
13 14 15 16	described in this subsection is a group health plan (other than a fully insured group health plan), that— "(1) provides coverage for obstetric care (such as pregnancy-related services) or routine gynecologic
 13 14 15 16 17 	described in this subsection is a group health plan (other than a fully insured group health plan), that— "(1) provides coverage for obstetric care (such as pregnancy-related services) or routine gynecologic care (such as preventive women's health examina-
 13 14 15 16 17 18 	described in this subsection is a group health plan (other than a fully insured group health plan), that— "(1) provides coverage for obstetric care (such as pregnancy-related services) or routine gynecologic care (such as preventive women's health examina- tions); and
 13 14 15 16 17 18 19 	described in this subsection is a group health plan (other than a fully insured group health plan), that— "(1) provides coverage for obstetric care (such as pregnancy-related services) or routine gynecologic care (such as preventive women's health examina- tions); and "(2) requires the designation by a participant or
 13 14 15 16 17 18 19 20 	described in this subsection is a group health plan (other than a fully insured group health plan), that— "(1) provides coverage for obstetric care (such as pregnancy-related services) or routine gynecologic care (such as preventive women's health examina- tions); and "(2) requires the designation by a participant or beneficiary of a participating primary care provider
 13 14 15 16 17 18 19 20 21 	described in this subsection is a group health plan (other than a fully insured group health plan), that— "(1) provides coverage for obstetric care (such as pregnancy-related services) or routine gynecologic care (such as preventive women's health examina- tions); and "(2) requires the designation by a participant or beneficiary of a participating primary care provider who is not a physician who specializes in obstetrics

1	"(1) as waiving any coverage requirement relat-
2	ing to medical necessity or appropriateness with re-
3	spect to the coverage of obstetric or gynecologic care
4	described in subsection (a);
5	"(2) to preclude the plan from requiring that the
6	physician who specializes in obstetrics or gynecology
7	notify the designated primary care provider or the
8	plan of treatment decisions;
9	"(3) to preclude a group health plan from allow-
10	ing health care professionals other than physicians to
11	provide routine obstetric or routine gynecologic care;
12	or
13	"(4) to preclude a group health plan from per-
14	mitting a physician who specializes in obstetrics and
15	gynecology from being a primary care provider under
16	the plan.
17	"SEC. 9824. PATIENT ACCESS TO PEDIATRIC CARE.
18	"(a) IN GENERAL.—In the case of a group health plan
19	(other than a fully insured group health plan) that provides
20	coverage for routine pediatric care and that requires the
21	designation by a participant or beneficiary of a partici-
22	pating primary care provider, if the designated primary
23	care provider is not a physician who specializes in
24	pediatrics—

1	"(1) the plan may not require authorization or
2	referral by the primary care provider in order for a
3	participant or beneficiary to obtain coverage for rou-
4	tine pediatric care; and
5	"(2) the plan shall treat the ordering of other
6	routine care related to routine pediatric care by such
7	a specialist as having been authorized by the des-
8	ignated primary care provider.
9	"(b) Rules of Construction.—Nothing in sub-
10	section (a) shall be construed—
11	"(1) as waiving any coverage requirement relat-
12	ing to medical necessity or appropriateness with re-
13	spect to the coverage of any pediatric care provided
14	to, or ordered for, a participant or beneficiary;
15	"(2) to preclude a group health plan from re-
16	quiring that a specialist described in subsection (a)
17	notify the designated primary care provider or the
18	plan of treatment decisions; or
19	"(3) to preclude a group health plan from allow-
20	ing health care professionals other than physicians to
21	provide routine pediatric care.
22	"SEC. 9825. TIMELY ACCESS TO SPECIALISTS.
23	"(a) TIMELY ACCESS.—
24	"(1) IN GENERAL.—A group health plan (other
25	than a fully insured group health plan) shall ensure

1	that participants and beneficiaries have timely, in
2	accordance with the medical exigencies of the case, ac-
3	cess to primary and specialty health care profes-
4	sionals who are appropriate to the condition of the
5	participant or beneficiary, when such care is covered
6	under the plan. Such access may be provided through
7	contractual arrangements with specialized providers
8	outside of the network of the plan.
9	"(2) RULE OF CONSTRUCTION.—Nothing in
10	paragraph (1) shall be construed—
11	(A) to require the coverage under a group
12	health plan of particular benefits or services or
13	to prohibit a plan from including providers only
14	to the extent necessary to meet the needs of the
15	plan's participants or beneficiaries or from es-
16	tablishing any measure designed to maintain
17	quality and control costs consistent with the re-
18	sponsibilities of the plan; or
19	"(B) to override any State licensure or
20	scope-of-practice law.
21	"(b) TREATMENT PLANS.—
22	"(1) IN GENERAL.—Nothing in this section shall
23	be construed to prohibit a group health plan (other
24	than a fully insured group health plan) from requir-

1	ing that specialty care be provided pursuant to a
2	treatment plan so long as the treatment plan is—
3	"(A) developed by the specialist, in con-
4	sultation with the case manager or primary care
5	provider, and the participant or beneficiary;
6	"(B) approved by the plan in a timely
7	manner in accordance with the medical exigen-
8	cies of the case; and
9	``(C) in accordance with the applicable
10	quality assurance and utilization review stand-
11	ards of the plan.
12	"(2) NOTIFICATION.—Nothing in paragraph (1)
13	shall be construed as prohibiting a plan from requir-
14	ing the specialist to provide the case manager or pri-
15	mary care provider with regular updates on the spe-
16	cialty care provided, as well as all other necessary
17	medical information.
18	"(c) Referrals.—Nothing in this section shall be
19	construed to prohibit a plan from requiring an authoriza-
20	tion by the case manager or primary care provider of the
21	participant or beneficiary in order to obtain coverage for
22	specialty services so long as such authorization is for an
23	adequate number of referrals.
24	"(d) Specialty Care Defined.—For purposes of this

subsection, the term 'specialty care' means, with respect to

a condition, care and treatment provided by a health care
 practitioner, facility, or center (such as a center of excel lence) that has adequate expertise (including age-appro priate expertise) through appropriate training and experi ence.

6 "SEC. 9826. CONTINUITY OF CARE.

7 "(a) IN GENERAL.—

"(1) TERMINATION OF PROVIDER.—If a contract 8 9 between a group health plan (other than a fully in-10 sured group health plan) and a health care provider 11 is terminated (as defined in paragraph (2)), or bene-12 fits or coverage provided by a health care provider are 13 terminated because of a change in the terms of pro-14 vider participation in such group health plan, and 15 an individual who is a participant or beneficiary in 16 the plan is undergoing a course of treatment from the 17 provider at the time of such termination, the plan 18 shall—

19	"(A) notify the individual on a timely basis
20	of such termination;

21 "(B) provide the individual with an oppor22 tunity to notify the plan of a need for transi23 tional care; and

24 "(C) in the case of termination described in
25 paragraph (2), (3), or (4) of subsection (b), and

subject to subsection (c), permit the individual to
continue or be covered with respect to the course
of treatment with the provider's consent during
a transitional period (as provided under sub-
section (b)).
"(2) TERMINATED.—In this section, the term
'terminated' includes, with respect to a contract, the
expiration or nonrenewal of the contract by the group
health plan, but does not include a termination of the
contract by the plan for failure to meet applicable
quality standards or for fraud.
"(3) CONTRACTS.—For purposes of this section,
the term 'contract between a group health plan (other
than a fully insured group health plan) and a health
care provider' shall include a contract between such
a plan and an organized network of providers.
"(b) Transitional Period.—
"(1) GENERAL RULE.—Except as provided in
paragraph (3), the transitional period under this sub-
section shall permit the participant or beneficiary to
extend the coverage involved for up to 90 days from
the date of the notice described in subsection $(a)(1)(A)$
of the provider's termination.
"(2) INSTITUTIONAL CARE.—Subject to para-
graph (1), the transitional period under this sub-

1	section for institutional or inpatient care from a pro-
2	vider shall extend until the discharge or termination
3	of the period of institutionalization and also shall in-
4	clude institutional care provided within a reasonable
5	time of the date of termination of the provider status
6	if the care was scheduled before the date of the an-
7	nouncement of the termination of the provider status
8	under subsection $(a)(1)(A)$ or if the individual on
9	such date was on an established waiting list or other-
10	wise scheduled to have such care.
11	"(3) Pregnancy.—Notwithstanding paragraph
12	(1), if—
13	``(A) a participant or beneficiary has en-
14	tered the second trimester of pregnancy at the
15	time of a provider's termination of participa-
16	tion; and
17	``(B) the provider was treating the preg-
18	nancy before the date of the termination;
19	the transitional period under this subsection with re-
20	spect to provider's treatment of the pregnancy shall
21	extend through the provision of post-partum care di-
22	rectly related to the delivery.
23	"(4) TERMINAL ILLNESS.—Notwithstanding
24	paragraph (1), if—

1	"(A) a participant or beneficiary was deter-
2	mined to be terminally ill (as determined under
3	section 1861(dd)(3)(A) of the Social Security
4	Act) prior to a provider's termination of partici-
5	pation; and
6	``(B) the provider was treating the terminal
7	illness before the date of termination;
8	the transitional period under this subsection shall be
9	for care directly related to the treatment of the ter-
10	minal illness and shall extend for the remainder of
11	the individual's life for such care.
12	"(c) Permissible Terms and Conditions.—A group
13	health plan (other than a fully insured group health plan)
14	may condition coverage of continued treatment by a pro-
15	vider under subsection $(a)(1)(C)$ upon the provider agreeing
16	to the following terms and conditions:
17	"(1) The provider agrees to accept reimburse-
18	ment from the plan and individual involved (with re-
19	spect to cost-sharing) at the rates applicable prior to
20	the start of the transitional period as payment in full
21	(or at the rates applicable under the replacement plan
22	after the date of the termination of the contract with
23	the group health plan) and not to impose cost-sharing
24	with respect to the individual in an amount that
25	would exceed the cost-sharing that could have been

1	imposed if the contract referred to in subsection (a)(1)
2	had not been terminated.
3	"(2) The provider agrees to adhere to the quality
4	assurance standards of the plan responsible for pay-
5	ment under paragraph (1) and to provide to such
6	plan necessary medical information related to the
7	care provided.
8	"(3) The provider agrees otherwise to adhere to
9	such plan's policies and procedures, including proce-
10	dures regarding referrals and obtaining prior author-
11	ization and providing services pursuant to a treat-
12	ment plan (if any) approved by the plan.
13	"(d) Rule of Construction.—Nothing in this sec-
14	tion shall be construed to require the coverage of benefits
15	which would not have been covered if the provider involved
16	remained a participating provider.
17	"(e) DEFINITION.—In this section, the term health
18	care provider' or 'provider' means—
19	"(1) any individual who is engaged in the deliv-
20	ery of health care services in a State and who is re-
21	quired by State law or regulation to be licensed or
22	certified by the State to engage in the delivery of such
23	services in the State; and
24	"(2) any entity that is engaged in the delivery

57

25 of health care services in a State and that, if it is re-

quired by State law or regulation to be licensed or
 certified by the State to engage in the delivery of such
 services in the State, is so licensed.

4 "(f) COMPREHENSIVE STUDY OF COST, QUALITY AND
5 COORDINATION OF COVERAGE FOR PATIENTS AT THE END
6 OF LIFE.—

7 "(1) Study by the medicare payment advi-SORY COMMISSION.—The Medicare Payment Advisory 8 9 Commission shall conduct a study of the costs and 10 patterns of care for persons with serious and complex 11 conditions and the possibilities of improving upon 12 that care to the degree it is triggered by the current 13 category of terminally ill as such term is used for 14 purposes of section 1861(dd) of the Social Security 15 Act (relating to hospice benefits) or of utilizing care 16 in other payment settings in Medicare.

17 "(2) AGENCY FOR HEALTH CARE POLICY AND RE18 SEARCH.—The Agency for Health Care Policy and
19 Research shall conduct studies of the possible thresh20 olds for major conditions causing serious and complex
21 illness, their administrative parameters and feasi22 bility, and their impact upon costs and quality.

23 "(3) HEALTH CARE FINANCING ADMINISTRA24 TION.—The Health Care Financing Administration
25 shall conduct studies of the merits of applying similar

59 thresholds in Medicare+Choice programs, including

thresholds in Medicare+Choice programs, including
 adapting risk adjustment methods to account for this
 category.

"(4) Initial report.—

4

"(A) IN GENERAL.—Not later than 12 5 6 months after the date of enactment of this sec-7 tion, the Medicare Payment Advisory Commis-8 sion and the Agency for Health Care Policy and 9 Research shall each prepare and submit to the 10 Committee on Health, Education, Labor and 11 Pensions of the Senate a report concerning the 12 results of the studies conducted under para-13 graphs (1) and (2), respectively.

14 "(B) COPY TO SECRETARY.—Concurrent
15 with the submission of the reports under sub16 paragraph (A), the Medicare Payment Advisory
17 Commission and the Agency for health Care Pol18 icy and Research shall transmit a copy of the re19 ports under such subparagraph to the Secretary.
20 "(5) FINAL REPORT.—

21 "(A) CONTRACT WITH INSTITUTE OF MEDI22 CINE.—Not later than 1 year after the submis23 sion of the reports under paragraph (4), the Sec24 retary of Health and Human Services shall con25 tract with the Institute of Medicine to conduct a

study of the practices and their effects arising
 from the utilization of the category "serious and
 complex" illness.

4 "(B) REPORT.—Not later than 1 year after
5 the date of the execution of the contract referred
6 to in subparagraph (A), the Institute of Medicine
7 shall prepare and submit to the Committee on
8 Health, Education, Labor and Pensions of the
9 Senate a report concerning the study conducted
10 pursuant to such contract.

"(6) FUNDING.—From funds appropriated to the
Department of Health and Human Services, the Secretary of Health and Human Services shall make
available such funds as the Secretary determines is
necessary to carry out this subsection.

16 "SEC. 9827. PROTECTION OF PATIENT-PROVIDER COMMU-

17

NICATIONS.

18 "(a) IN GENERAL.—Subject to subsection (b), a group health plan (other than a fully insured group health plan 19 and in relation to a participant or beneficiary) shall not 20 21 prohibit or otherwise restrict a health care professional from 22 advising such a participant or beneficiary who is a patient 23 of the professional about the health status of the participant 24 or beneficiary or medical care or treatment for the condition or disease of the participant or beneficiary, regardless of 25

whether coverage for such care or treatment are provided
 under the contract, if the professional is acting within the
 lawful scope of practice.

4 "(b) RULE OF CONSTRUCTION.—Nothing in this sec5 tion shall be construed as requiring a group health plan
6 (other than a fully insured group health plan) to provide
7 specific benefits under the terms of such plan.

8 "SEC. 9828. PATIENT'S RIGHT TO PRESCRIPTION DRUGS.

9 "To the extent that a group health plan (other than 10 a fully insured group health plan) provides coverage for 11 benefits with respect to prescription drugs, and limits such 12 coverage to drugs included in a formulary, the plan shall—

"(1) ensure the participation of physicians and
pharmacists in developing and reviewing such formulary; and

"(2) in accordance with the applicable quality
assurance and utilization review standards of the
plan, provide for exceptions from the formulary limitation when a non-formulary alternative is medically
necessary and appropriate.

21 "SEC. 9829. SELF-PAYMENT FOR BEHAVIORAL HEALTH CARE
22 SERVICES.

23 "(a) IN GENERAL.—A group health plan (other than
24 a fully insured group health plan) may not—

1	"(1) prohibit or otherwise discourage a partici-
2	pant or beneficiary from self-paying for behavioral
3	health care services once the plan has denied coverage
4	for such services; or
5	"(2) terminate a health care provider because
6	such provider permits participants or beneficiaries to
7	self-pay for behavioral health care services—
8	"(A) that are not otherwise covered under
9	the plan; or
10	"(B) for which the group health plan pro-
11	vides limited coverage, to the extent that the
12	group health plan denies coverage of the services.
13	"(b) RULE OF CONSTRUCTION.—Nothing in subsection
14	(a)(2)(B) shall be construed as prohibiting a group health
15	plan from terminating a contract with a health care pro-
16	vider for failure to meet applicable quality standards or
17	for fraud.
18	"SEC. 9830. COVERAGE FOR INDIVIDUALS PARTICIPATING
19	IN APPROVED CANCER CLINICAL TRIALS.
20	"(a) Coverage.—
21	"(1) IN GENERAL.—If a group health plan (other
22	than a fully insured group health plan) provides cov-
23	erage to a qualified individual (as defined in sub-
24	section (b)), the plan—

1	"(A) may not deny the individual partici-
2	pation in the clinical trial referred to in sub-
3	section $(b)(2);$
4	"(B) subject to subsections (b), (c), and (d)
5	may not deny (or limit or impose additional
6	conditions on) the coverage of routine patient
7	costs for items and services furnished in connec-
8	tion with participation in the trial; and
9	"(C) may not discriminate against the in-
10	dividual on the basis of the participant's or
11	beneficiaries participation in such trial.
12	"(2) Exclusion of certain costs.—For pur-
13	poses of paragraph $(1)(B)$, routine patient costs do
14	not include the cost of the tests or measurements con-
15	ducted primarily for the purpose of the clinical trial
16	involved.
17	"(3) Use of in-network providers.—If one or
18	more participating providers is participating in a
19	clinical trial, nothing in paragraph (1) shall be con-
20	strued as preventing a plan from requiring that a
21	qualified individual participate in the trial through
22	such a participating provider if the provider will ac-

24 "(b) QUALIFIED INDIVIDUAL DEFINED.—For purposes
25 of subsection (a), the term "qualified individual" means an

cept the individual as a participant in the trial.

23

1	individual who is a participant or beneficiary in a group
2	health plan and who meets the following conditions:
3	((1)(A) The individual has been diagnosed with
4	cancer for which no standard treatment is effective.
5	``(B) The individual is eligible to participate in
6	an approved clinical trial according to the trial pro-
7	tocol with respect to treatment of such illness.
8	``(C) The individual's participation in the trial
9	offers meaningful potential for significant clinical
10	benefit for the individual.
11	"(2) <i>Either</i> —
12	"(A) the referring physician is a partici-
13	pating health care professional and has con-
14	cluded that the individual's participation in
15	such trial would be appropriate based upon the
16	individual meeting the conditions described in
17	paragraph (1); or
18	"(B) the participant or beneficiary provides
19	medical and scientific information establishing
20	that the individual's participation in such trial
21	would be appropriate based upon the individual
22	meeting the conditions described in paragraph
23	(1).
24	"(c) PAYMENT.—

1	"(1) IN GENERAL.—Under this section a group
2	health plan (other than a fully insured group health
3	plan) shall provide for payment for routine patient
4	costs described in subsection $(a)(2)$ but is not required
5	to pay for costs of items and services that are reason-
6	ably expected to be paid for by the sponsors of an ap-
7	proved clinical trial.
8	"(2) Standards for determining routine
9	PATIENT COSTS ASSOCIATED WITH CLINICAL TRIAL
10	PARTICIPATION.—
11	"(A) IN GENERAL.—The Secretary shall es-
12	tablish, on an expedited basis and using a nego-
13	tiated rulemaking process under subchapter III
14	of chapter 5 of title 5, United States Code, stand-
15	ards relating to the coverage of routine patient
16	costs for individuals participating in clinical
17	trials that group health plans must meet under
18	this section.
19	"(B) FACTORS.—In establishing routine pa-
20	tient cost standards under subparagraph (A) , the
21	Secretary shall consult with interested parties
22	and take into account —
23	"(i) quality of patient care;
24	"(ii) routine patient care costs versus
25	costs associated with the conduct of clinical

1	trials, including unanticipated patient care
2	costs as a result of participation in clinical
3	trials; and
4	"(iii) previous and on-going studies re-
5	lating to patient care costs associated with
6	participation in clinical trials.
7	"(C) PUBLICATION OF NOTICE.—In car-
8	rying out the rulemaking process under this
9	paragraph, the Secretary, after consultation with
10	organizations representing cancer patients,
11	health care practitioners, medical researchers,
12	employers, group health plans, manufacturers of
13	drugs, biologics and medical devices, medical
14	economists, hospitals, and other interested par-
15	ties, shall publish notice provided for under sec-
16	tion 564(a) of title 5, United States Code, by not
17	later than 45 days after the date of the enact-
18	ment of this section.
19	(D) Target date for publication of
20	RULE.—As part of the notice under subpara-
21	graph (C), and for purposes of this paragraph,
22	the 'target date for publication' (referred to in
23	section $564(a)(5)$ of such title 5) shall be June
24	20, 2000

24 30, 2000.

66

1	"(E) Abbreviated period for submis-
2	SION OF COMMENTS.—In applying section 564(c)
3	of such title 5 under this paragraph, '15 days'
4	shall be substituted for '30 days'.
5	"(F) Appointment of negotiated rule-
6	MAKING COMMITTEE AND FACILITATOR.—The
7	Secretary shall provide for—
8	"(i) the appointment of a negotiated
9	rule making committee under section 565(a)
10	of such title 5 by not later than 30 days
11	after the end of the comment period pro-
12	vided for under section $564(c)$ of such title
13	5 (as shortened under subparagraph (E)),
14	and
15	"(ii) the nomination of a facilitator
16	under section 566(c) of such title 5 by not
17	later than 10 days after the date of appoint-
18	ment of the committee.
19	"(G) Preliminary committee report.—
20	The negotiated rulemaking committee appointed
21	under subparagraph (F) shall report to the Sec-
22	retary, by not later than March 29, 2000, re-
23	garding the committee's progress on achieving a
24	consensus with regard to the rulemaking pro-
25	ceeding and whether such consensus is likely to

1	occur before 1 month before the target date for
2	publication of the rule. If the committee reports
3	that the committee has failed to make significant
4	progress towards such consensus or is unlikely to
5	reach such consensus by the target date, the Sec-
6	retary may terminate such process and provide
7	for the publication of a rule under this para-
8	graph through such other methods as the Sec-
9	retary may provide.
10	"(H) FINAL COMMITTEE REPORT.—If the
11	committee is not terminated under subparagraph
12	(G), the rulemaking committee shall submit a re-
13	port containing a proposed rule by not later
14	than 1 month before the target date of publica-
15	tion.
16	"(I) FINAL EFFECT.—The Secretary shall
17	publish a rule under this paragraph in the Fed-
18	eral Register by not later than the target date of
19	publication.
20	"(J) Publication of rule after public
21	COMMENT.—The Secretary shall provide for con-
22	sideration of such comments and republication of
23	such rule by not later than 1 year after the tar-
24	get date of publication.

1	"(K) EFFECTIVE DATE.—The provisions of			
2	this paragraph shall apply to group health plans			
3	(other than a fully insured group health plan)			
4	for plan years beginning on or after January 1,			
5	2001.			
6	"(3) PAYMENT RATE.—In the case of covered			
7	items and services provided by—			
8	"(A) a participating provider, the payment			
9	rate shall be at the agreed upon rate, or			
10	``(B) a nonparticipating provider, the pay-			
11	ment rate shall be at the rate the plan would			
12	normally pay for comparable services under sub-			
13	paragraph (A).			
14	"(d) Approved Clinical Trial Defined.—			
15	"(1) IN GENERAL.—In this section, the term 'ap-			
16	proved clinical trial' means a cancer clinical research			
17	study or cancer clinical investigation approved and			
18	funded (which may include funding through in-kind			
19	contributions) by one or more of the following:			
20	"(A) The National Institutes of Health.			
21	(B) A cooperative group or center of the			
22	National Institutes of Health.			
23	"(C) Either of the following if the condi-			
24	tions described in paragraph (2) are met:			

10						
"(i) The Department of Veterans Af-						
fairs.						
"(ii) The Department of Defense.						
"(2) Conditions for departments.—The con-						
ditions described in this paragraph, for a study or in-						
vestigation conducted by a Department, are that the						
study or investigation has been reviewed and ap-						
proved through a system of peer review that the Sec-						
retary determines—						
"(A) to be comparable to the system of peer						
1 review of studies and investigations used by th						
National Institutes of Health, and						
(B) assures unbiased review of the highest						
scientific standards by qualified individuals who						
have no interest in the outcome of the review.						
"(e) CONSTRUCTION.—Nothing in this section shall be						
construed to limit a plan's coverage with respect to clinical						
trials.						
"(f) Plan Satisfaction of Certain Requirements;						
Responsibilities of Fiduciaries.—						
"(1) IN GENERAL.—For purposes of this section,						
insofar as a group health plan provides benefits in the						
form of health insurance coverage through a health in-						
surance issuer, the plan shall be treated as meeting						
the requirements of this section with respect to such						

1	benefits and not be considered as failing to meet such
2	requirements because of a failure of the issuer to meet
3	such requirements so long as the plan sponsor or its
4	representatives did not cause such failure by the
5	issuer.
6	"(2) CONSTRUCTION.—Nothing in this section
7	shall be construed to affect or modify the responsibil-
8	ities of the fiduciaries of a group health plan under
9	part 4 of subtitle B of title I of the Employee Retire-
10	ment Income Security Act of 1974.
11	"(g) Study and Report.—
12	"(1) Study.—The Secretary shall study the im-
13	pact on group health plans for covering routine pa-
14	tient care costs for individuals who are entitled to
15	benefits under this section and who are enrolled in an
16	approved cancer clinical trial program.
17	"(2) Report to congress.—Not later than
18	January 1, 2005, the Secretary shall submit a report
19	to Congress that contains an assessment of—
20	"(A) any incremental cost to group health
21	plans resulting from the provisions of this sec-
22	tion;
23	(B) a projection of expenditures to such
24	plans resulting from this section; and

1	"(C)	any	impact	on	premiums	resulting
2	from this s	section	n.			

3 "SEC. 9830A. PROHIBITING DISCRIMINATION AGAINST PRO4 VIDERS.

"(a) IN GENERAL.—A group health plan (other than 5 a fully insured group health plan) shall not discriminate 6 7 with respect to participation or indemnification as to any 8 provider who is acting within the scope of the provider's 9 license or certification under applicable State law, solely 10 on the basis of such license or certification. This subsection shall not be construed as requiring the coverage under a 11 plan of particular benefits or services or to prohibit a plan 12 13 from including providers only to the extent necessary to meet the needs of the plan's participants and beneficiaries 14 15 or from establishing any measure designed to maintain quality and control costs consistent with the responsibilities 16 17 of the plan.

18 "(b) NO REQUIREMENT FOR ANY WILLING PRO19 VIDER.—Nothing in this section shall be construed as re20 quiring a group health plan that offers network coverage
21 to include for participation every willing provider or health
22 professional who meets the terms and conditions of the plan.
23 "SEC. 9830B. GENERALLY APPLICABLE PROVISION.

24 "In the case of a group health plan that provides bene-25 fits under 2 or more coverage options, the requirements of
this subchapter shall apply separately with respect to each
 coverage option.".

3 (b) DEFINITION.—Section 9832(b) of the Internal Rev4 enue Code of 1986 is amended by adding at the end the
5 following:

6 "(4) FULLY INSURED GROUP HEALTH PLAN.— 7 The term 'fully insured group health plan' means a 8 group health plan where benefits under the plan are 9 provided pursuant to the terms of an arrangement be-10 tween a group health plan and a health insurance 11 issuer and are guaranteed by the health insurance 12 issuer under a contract or policy of insurance.".

(c) CONFORMING AMENDMENT.—Chapter 98 of the Internal Revenue Code of 1986 is amended in the table of subchapters in the item relating to subchapter C, by striking
"Subchapter C" and inserting "Subchapter D".

17 SEC. 103. EFFECTIVE DATE AND RELATED RULES.

(a) IN GENERAL.—The amendments made by this subtitle shall apply with respect to plan years beginning on
or after January 1 of the second calendar year following
the date of the enactment of this Act. The Secretary shall
issue all regulations necessary to carry out the amendments
made by this section before the effective date thereof.

(b) LIMITATION ON ENFORCEMENT ACTIONS.—No en25 forcement action shall be taken, pursuant to the amend-

ments made by this subtitle, against a group health plan
 with respect to a violation of a requirement imposed by such
 amendments before the date of issuance of regulations issued
 in connection with such requirement, if the plan has sought
 to comply in good faith with such requirement.

6 Subtitle B—Right to Information 7 About Plans and Providers

8 SEC. 111. INFORMATION ABOUT PLANS.

9 (a) EMPLOYEE RETIREMENT INCOME SECURITY ACT 10 OF 1974.—

(1) IN GENERAL.—Subpart B of part 7 of subtitle B of title I of the Employee Retirement Income
Security Act of 1974 (29 U.S.C. 1185 et seq.) is
amended by adding at the end the following:

15 "SEC. 714. HEALTH PLAN COMPARATIVE INFORMATION.

16 "(a) REQUIREMENT.—

17 "(1) IN GENERAL.—A group health plan, and a 18 health insurance issuer that provides coverage in con-19 nection with group health insurance coverage, shall, 20 not later than 12 months after the date of enactment 21 of this section, and at least annually thereafter, pro-22 vide for the disclosure, in a clear and accurate form 23 to each participant and each beneficiary who does not 24 reside at the same address as the participant, or upon 25 request to an individual eligible for coverage under the plan, of the information described in subsection
 (b).

3 "(2) RULE OF CONSTRUCTION.—Nothing in this 4 section shall be construed to prevent a plan or issuer 5 from entering into any agreement under which the 6 issuer agrees to assume responsibility for compliance 7 with the requirements of this section and the plan is 8 released from liability for such compliance. 9 "(3) PROVISION OF INFORMATION.—Information 10 shall be provided to participants and beneficiaries 11 under this section at the address maintained by the

12 plan or issuer with respect to such participants or13 beneficiaries.

14 "(b) REQUIRED INFORMATION.—The informational
15 materials to be distributed under this section shall include
16 for each package option available under a group health plan
17 the following:

"(1) A description of the covered items and services under each such plan and any in- and out-of-network features of each such plan, including a summary description of the specific exclusions from coverage under the plan.

23 "(2) A description of any cost-sharing, including
24 premiums, deductibles, coinsurance, and copayment
25 amounts, for which the participant or beneficiary will

be responsible, including any annual or lifetime lim-
its on benefits, for each such plan.
"(3) A description of any optional supplemental
benefits offered by each such plan and the terms and
conditions (including premiums or cost-sharing) for
such supplemental coverage.
"(4) A description of any restrictions on pay-
ments for services furnished to a participant or bene-
ficiary by a health care professional that is not a
participating professional and the liability of the
participant or beneficiary for additional payments
for these services.
"(5) A description of the service area of each
such plan, including the provision of any out-of-area
coverage.
"(6) A description of the extent to which partici-
pants and beneficiaries may select the primary care
provider of their choice, including providers both
within the network and outside the network of each
such plan (if the plan permits out-of-network serv-
ices).
"(7) A description of the procedures for advance
directives and organ donation decisions if the plan
maintains such procedures.

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1	"(8) A description of the requirements and pro-
2	cedures to be used to obtain preauthorization for
3	health services (including telephone numbers and
4	mailing addresses), including referrals for specialty
5	care.
6	"(9) A description of the definition of medical
7	necessity used in making coverage determinations by
8	each such plan.
9	"(10) A summary of the rules and methods for
10	appealing coverage decisions and filing grievances
11	(including telephone numbers and mailing addresses),
12	as well as other available remedies.
13	"(11) A summary description of any provisions
14	for obtaining off-formulary medications if the plan
15	utilizes a defined formulary for providing specific
16	prescription medications.
17	"(12) A summary of the rules for access to emer-
18	gency room care. Also, any available educational ma-
19	terial regarding proper use of emergency services.
20	"(13) A description of whether or not coverage is
21	provided for experimental treatments, investigational
22	treatments, or clinical trials and the circumstances
23	under which access to such treatments or trials is
24	made available.

1	"(14) A description of the specific preventative
2	services covered under the plan if such services are
3	covered.
4	"(15) A statement regarding—
5	"(A) the manner in which a participant or
6	beneficiary may access an obstetrician, gyne-
7	cologist, or pediatrician in accordance with sec-
8	tion 723 or 724; and
9	(B) the manner in which a participant or
10	beneficiary obtains continuity of care as pro-
11	vided for in section 726.
12	((16) A statement that the following informa-
13	tion, and instructions on obtaining such information
14	(including telephone numbers and, if available, Inter-
15	net websites), shall be made available upon request:
16	"(A) The names, addresses, telephone num-
17	bers, and State licensure status of the plan's par-
18	ticipating health care professionals and partici-
19	pating health care facilities, and, if available,
20	the education, training, specialty qualifications
21	or certifications of such professionals.
22	(B) A summary description of the methods
23	used for compensating participating health care
24	professionals, such as capitation, fee-for-service,
25	salary, or a combination thereof. The require-

1	ment of this subparagraph shall not be construed
2	as requiring plans to provide information con-
3	cerning proprietary payment methodology.
4	"(C) A summary description of the methods
5	used for compensating health care facilities, in-
6	cluding per diem, fee-for-service, capitation, bun-
7	dled payments, or a combination thereof. The re-
8	quirement of this subparagraph shall not be con-
9	strued as requiring plans to provide information
10	concerning proprietary payment methodology.
11	(D) A summary description of the proce-
12	dures used for utilization review.
13	"(E) The list of the specific prescription
14	medications included in the formulary of the
15	plan, if the plan uses a defined formulary.
16	"(F) A description of the specific exclusions
17	from coverage under the plan.
18	``(G) Any available information related to
19	the availability of translation or interpretation
20	services for non-English speakers and people
21	with communication disabilities, including the
22	availability of audio tapes or information in
23	Braille.
24	"(H) Any information that is made public
25	by accrediting organizations in the process of ac-

creditation if the plan is accredited, or any ad ditional quality indicators that the plan makes
 available.

4 "(c) MANNER OF DISTRIBUTION.—The information de5 scribed in this section shall be distributed in an accessible
6 format that is understandable to an average plan partici7 pant or beneficiary.

8 "(d) RULE OF CONSTRUCTION.—Nothing in this sec-9 tion may be construed to prohibit a group health plan, or health insurance issuer in connection with group health in-10 surance coverage, from distributing any other additional 11 information determined by the plan or issuer to be impor-12 tant or necessary in assisting participants and beneficiaries 13 or upon request potential participants and beneficiaries in 14 15 the selection of a health plan or from providing information under subsection (b)(15) as part of the required informa-16 tion. 17

18 "(e) CONFORMING REGULATIONS.—The Secretary 19 shall issue regulations to coordinate the requirements on 20 group health plans and health insurance issuers under this 21 section with the requirements imposed under part 1, to re-22 duce duplication with respect to any information that is 23 required to be provided under any such requirements.

24 "(f) HEALTH CARE PROFESSIONAL.—In this section,
25 the term 'health care professional' means a physician (as

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defined in section 1861(r) of the Social Security Act) or 1 2 other health care professional if coverage for the professional's services is provided under the health plan involved 3 for the services of the professional. Such term includes a 4 5 podiatrist, optometrist, chiropractor, psychologist, dentist, physician assistant, physical or occupational therapist and 6 7 therapy assistant, speech-language pathologist, audiologist, 8 registered or licensed practical nurse (including nurse prac-9 titioner, clinical nurse specialist, certified registered nurse anesthetist, and certified nurse-midwife), licensed certified 10 11 social worker, registered respiratory therapist, and certified 12 respiratory therapy technician.".

13 (2) Conforming Amendments.—

- 14 (A) Section 732(a) of the Employee Retire15 ment Income Security Act of 1974 (29 U.S.C.
 16 1191a(a)) is amended by striking "section 711,
 17 and inserting "sections 711 and 714".
 18 (B) The table of contents in section 1 of the
- 19Employee Retirement Income Security Act of201974 (29 U.S.C. 1001) is amended by inserting21after the item relating to section 713, the fol-22lowing:

"Sec. 714. Health plan comparative information.".

23 (b) INTERNAL REVENUE CODE OF 1986.—Subchapter
24 B of chapter 100 of the Internal Revenue Code of 1986 is
25 amended—

1	(1) in the table of sections, by inserting after the
2	item relating to section 9812 the following new item:
	"Sec. 9813. Health plan comparative information.";
3	and
4	(2) by inserting after section 9812 the following:
5	"SEC. 9813. HEALTH PLAN COMPARATIVE INFORMATION.
6	"(a) Requirement.—
7	"(1) IN GENERAL.—A group health plan shall,
8	not later than 12 months after the date of enactment
9	of this section, and at least annually thereafter, pro-
10	vide for the disclosure, in a clear and accurate form
11	to each participant and each beneficiary who does not
12	reside at the same address as the participant, or upon
13	request to an individual eligible for coverage under
14	the plan, of the information described in subsection
15	(b).
16	"(2) Rules of construction.—Nothing in this

16 "(2) RULES OF CONSTRUCTION.—Nothing in this 17 section shall be construed to prevent a plan from en-18 tering into any agreement under which a health in-19 surance issuer agrees to assume responsibility for 20 compliance with the requirements of this section and 21 the plan is released from liability for such compli-22 ance.

23 "(3) PROVISION OF INFORMATION.—Information
24 shall be provided to participants and beneficiaries
25 under this section at the address maintained by the
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plan with respect to such participants or bene ficiaries.

3 "(b) REQUIRED INFORMATION.—The informational
4 materials to be distributed under this section shall include
5 for each package option available under a group health plan
6 the following:

7 "(1) A description of the covered items and serv8 ices under each such plan and any in- and out-of-net9 work features of each such plan, including a sum10 mary description of the specific exclusions from cov11 erage under the plan.

"(2) A description of any cost-sharing, including
premiums, deductibles, coinsurance, and copayment
amounts, for which the participant or beneficiary will
be responsible, including any annual or lifetime limits on benefits, for each such plan.

17 "(3) A description of any optional supplemental
18 benefits offered by each such plan and the terms and
19 conditions (including premiums or cost-sharing) for
20 such supplemental coverage.

21 "(4) A description of any restrictions on pay22 ments for services furnished to a participant or bene23 ficiary by a health care professional that is not a
24 participating professional and the liability of the

participant or beneficiary for additional payments
 for these services.

3 "(5) A description of the service area of each
4 such plan, including the provision of any out-of-area
5 coverage.

6 "(6) A description of the extent to which partici-7 pants and beneficiaries may select the primary care 8 provider of their choice, including providers both 9 within the network and outside the network of each 10 such plan (if the plan permits out-of-network serv-11 ices).

12 "(7) A description of the procedures for advance
13 directives and organ donation decisions if the plan
14 maintains such procedures.

"(8) A description of the requirements and procedures to be used to obtain preauthorization for
health services (including telephone numbers and
mailing addresses), including referrals for specialty
care.

20 "(9) A description of the definition of medical
21 necessity used in making coverage determinations by
22 each such plan.

23 "(10) A summary of the rules and methods for
24 appealing coverage decisions and filing grievances

1	(including telephone numbers and mailing addresses),
2	as well as other available remedies.
3	"(11) A summary description of any provisions
4	for obtaining off-formulary medications if the plan
5	utilizes a defined formulary for providing specific
6	prescription medications.
7	"(12) A summary of the rules for access to emer-
8	gency room care. Also, any available educational ma-
9	terial regarding proper use of emergency services.
10	"(13) A description of whether or not coverage is
11	provided for experimental treatments, investigational
12	treatments, or clinical trials and the circumstances
13	under which access to such treatments or trials is
14	made available.
15	"(14) A description of the specific preventative
16	services covered under the plan if such services are
17	covered.
18	"(15) A statement regarding—
19	"(A) the manner in which a participant or
20	beneficiary may access an obstetrician, gyne-
21	cologist, or pediatrician in accordance with sec-
22	tion 723 or 724; and
23	(B) the manner in which a participant or
24	beneficiary obtains continuity of care as pro-
25	vided for in section 726.

1	((16) A statement that the following informa-
2	tion, and instructions on obtaining such information
3	(including telephone numbers and, if available, Inter-
4	net websites), shall be made available upon request:
5	"(A) The names, addresses, telephone num-
6	bers, and State licensure status of the plan's par-
7	ticipating health care professionals and partici-
8	pating health care facilities, and, if available,
9	the education, training, specialty qualifications
10	or certifications of such professionals.
11	(B) A summary description of the methods
12	used for compensating participating health care
13	professionals, such as capitation, fee-for-service,
14	salary, or a combination thereof. The require-
15	ment of this subparagraph shall not be construed
16	as requiring plans to provide information con-
17	cerning proprietary payment methodology.
18	"(C) A summary description of the methods
19	used for compensating health care facilities, in-
20	cluding per diem, fee-for-service, capitation, bun-
21	dled payments, or a combination thereof. The re-
22	quirement of this subparagraph shall not be con-
23	strued as requiring plans to provide information
24	concerning proprietary payment methodology.

1	(D) A summary description of the proce-
2	dures used for utilization review.
3	``(E) The list of the specific prescription
4	medications included in the formulary of the
5	plan, if the plan uses a defined formulary.
6	``(F) A description of the specific exclusions
7	from coverage under the plan.
8	"(G) Any available information related to
9	the availability of translation or interpretation
10	services for non-English speakers and people
11	with communication disabilities, including the
12	availability of audio tapes or information in
13	Braille.
14	``(H) Any information that is made public
15	by accrediting organizations in the process of ac-
16	creditation if the plan is accredited, or any ad-
17	ditional quality indicators that the plan makes
18	available.
19	"(c) MANNER OF DISTRIBUTION.—The information de-
20	scribed in this section shall be distributed in an accessible
21	format that is understandable to an average plan partici-
22	pant or beneficiary.
23	"(d) Rule of Construction.—Nothing in this sec-
24	tion may be construed to prohibit a group health plan from
25	distributing any other additional information determined

by the plan to be important or necessary in assisting par ticipants and beneficiaries or upon request potential par ticipants and beneficiaries in the selection of a health plan
 or from providing information under subsection (b)(15) as
 part of the required information.

6 "(e) HEALTH CARE PROFESSIONAL.—In this section, 7 the term 'health care professional' means a physician (as 8 defined in section 1861(r) of the Social Security Act) or 9 other health care professional if coverage for the profes-10 sional's services is provided under the health plan involved for the services of the professional. Such term includes a 11 podiatrist, optometrist, chiropractor, psychologist, dentist, 12 13 physician assistant, physical or occupational therapist and therapy assistant, speech-language pathologist, audiologist, 14 15 registered or licensed practical nurse (including nurse practitioner, clinical nurse specialist, certified registered nurse 16 anesthetist, and certified nurse-midwife), licensed certified 17 18 social worker, registered respiratory therapist, and certified 19 respiratory therapy technician.".

20 SEC. 112. INFORMATION ABOUT PROVIDERS.

(a) STUDY.—The Secretary of Health and Human
Services shall enter into a contract with the Institute of
Medicine for the conduct of a study, and the submission
to the Secretary of a report, that includes—

1	(1) an analysis of information concerning health
2	care professionals that is currently available to pa-
3	tients, consumers, States, and professional societies,
4	nationally and on a State-by-State basis, including
5	patient preferences with respect to information about
6	such professionals and their competencies;
7	(2) an evaluation of the legal and other barriers
8	to the sharing of information concerning health care
9	professionals; and
10	(3) recommendations for the disclosure of infor-
11	mation on health care professionals, including the
12	competencies and professional qualifications of such
13	practitioners, to better facilitate patient choice, qual-
14	ity improvement, and market competition.
15	(b) REPORT.—Not later than 18 months after the date
16	of enactment of this Act, the Secretary of Health and
17	Human Services shall forward to the appropriate commit-
18	tees of Congress a copy of the report and study conducted
19	under subsection (a).

Subtitle C—Right to Hold Health 1 **Plans** Accountable 2 3 SEC. 121. AMENDMENT TO EMPLOYEE RETIREMENT IN-4 COME SECURITY ACT OF 1974. 5 (a) IN GENERAL.—Section 503 of the Employee Retirement Income Security Act of 1974 (29 U.S.C. 1133) is 6 amended to read as follows: 7 8 "SEC. 503. CLAIMS PROCEDURE, COVERAGE DETERMINA-9 TION. GRIEVANCES AND APPEALS. 10 "(a) CLAIMS PROCEDURE.—In accordance with requ-11 lations of the Secretary, every employee benefit plan shall— 12 "(1) provide adequate notice in writing to any 13 participant or beneficiary whose claim for benefits 14 under the plan has been denied, setting forth the spe-15 cific reasons for such denial, written in a manner cal-16 culated to be understood by the participant; and 17 "(2) afford a reasonable opportunity to any par-18 ticipant whose claim for benefits has been denied for 19 a full and fair review by the appropriate named fidu-20 ciary of the decision denying the claim. 21 "(b) Coverage Determinations Under Group 22 HEALTH PLANS.— 23 "(1) Procedures.— 24 "(A) IN GENERAL.—A group health plan or

25 *health insurance issuer conducting utilization re-*

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view	shall	ensure	that	procedures	are	in	place
for—							

3	"(i) making determinations regarding
4	whether a participant or beneficiary is eli-
5	gible to receive a payment or coverage for
6	health services under the plan or coverage
7	involved and any cost-sharing amount that
8	the participant or beneficiary is required to
9	pay with respect to such service;

10 "(*ii*) notifying a covered participant or 11 beneficiary (or the authorized representative of such participant or beneficiary) and the 12 13 treating health care professionals involved 14 regarding determinations made under the 15 plan or issuer and any additional payments that the participant or beneficiary 16 17 may be required to make with respect to 18 such service; and

"(iii) responding to requests, either
written or oral, for coverage determinations
or for internal appeals from a participant
or beneficiary (or the authorized representative of such participant or beneficiary) or
the treating health care professional with

1	the consent of the participant or bene-
2	ficiary.
3	"(B) ORAL REQUESTS.—With respect to an
4	oral request described in subparagraph (A)(iii),
5	a group health plan or health insurance issuer
6	may require that the requesting individual pro-
7	vide written evidence of such request.
8	"(2) Timeline for making determinations.—
9	"(A) ROUTINE DETERMINATION.—A group
10	health plan or a health insurance issuer shall
11	maintain procedures to ensure that prior author-
12	ization determinations concerning the provision
13	of non-emergency items or services are made
14	within 30 days from the date on which the re-
15	quest for a determination is submitted, except
16	that such period may be extended where certain
17	circumstances exist that are determined by the
18	Secretary to be beyond control of the plan or
19	issuer.
20	"(B) Expedited determination.—
21	"(i) In General.—A prior authoriza-

2 22 tion determination under this subsection 23 shall be made within 72 hours, in accord-24 ance with the medical exigencies of the case,

1	after a request is received by the plan or
2	issuer under clause (ii) or (iii).
3	"(ii) Request by participant or
4	BENEFICIARY.—A plan or issuer shall
5	maintain procedures for expediting a prior
6	authorization determination under this sub-
7	section upon the request of a participant or
8	beneficiary if, based on such a request, the
9	plan or issuer determines that the normal
10	time for making such a determination could
11	seriously jeopardize the life or health of the
12	participant or beneficiary.
13	"(iii) Documentation by health
14	CARE PROFESSIONAL.—A plan or issuer
15	shall maintain procedures for expediting a
16	prior authorization determination under
17	this subsection if the request involved indi-
18	cates that the treating health care profes-
19	sional has reasonably documented, based on
20	the medical exigencies, that a determination
21	under the procedures described in subpara-
22	graph (A) could seriously jeopardize the life
23	or health of the participant or beneficiary.
24	"(C) Concurrent determinations.—A
25	plan or issuer shall maintain procedures to cer-

1	tify or deny coverage of an extended stay or ad-
2	ditional services.
3	"(D) Retrospective determination.—A
4	plan or issuer shall maintain procedures to en-
5	sure that, with respect to the retrospective review
6	of a determination made under paragraph (1),
7	the determination shall be made within 30 work-
8	ing days of the date on which the plan or issuer
9	receives necessary information.
10	"(3) Notice of determinations.—
11	"(A) ROUTINE DETERMINATION.—With re-
12	spect to a coverage determination of a plan or
13	issuer under paragraph (2)(A), the plan or
14	issuer shall issue notice of such determination to
15	the participant or beneficiary (or the authorized
16	representative of the participant or beneficiary)
17	and, consistent with the medical exigencies of the
18	case, to the treating health care professional in-
19	volved not later than 2 working days after the
20	date on which the determination is made.
21	"(B) Expedited determination.—With
22	respect to a coverage determination of a plan or
23	issuer under paragraph (2)(B), the plan or
24	issuer shall issue notice of such determination to

1	representative of the participant or beneficiary),
2	and consistent with the medical exigencies of the
3	case, to the treating health care professional in-
4	volved within the 72 hour period described in
5	paragraph (2)(B).
6	"(C) Concurrent reviews.—With respect
7	to the determination under a plan or issuer
8	under paragraph $(2)(C)$ to certify or deny cov-
9	erage of an extended stay or additional services,
10	the plan or issuer shall issue notice of such deter-
11	mination to the treating health care professional
12	and to the participant or beneficiary involved
13	(or the authorized representative of the partici-
14	pant or beneficiary) within 1 working day of the
15	determination.
16	"(D) Retrospective reviews.—With re-
17	spect to the retrospective review under a plan or
18	issuer of a determination made under paragraph
19	(2)(D), the plan or issuer shall issue written no-
20	tice of an approval or disapproval of a deter-
21	mination under this subparagraph to the partic-
22	ipant or beneficiary (or the authorized represent-
23	ative of the participant or beneficiary) and
24	health care provider involved within 5 working

days of the date on which such determination is made.

"(E) REQUIREMENTS OF NOTICE OF AD-3 4 VERSE COVERAGE DETERMINATIONS.—A written notice of an adverse coverage determination 5 6 under this subsection, or of an expedited adverse 7 coverage determination under paragraph (2)(B), 8 shall be provided to the participant or bene-9 ficiary (or the authorized representative of the 10 participant or beneficiary) and treating health 11 care professional (if any) involved and shall 12 include— 13 "(i) the reasons for the determination 14 (including the clinical or scientific-evidence 15 based rationale used in making the deter-16 mination) written in a manner to be under-17 standable to the average participant or ben-18 eficiary; 19 "(*ii*) the procedures for obtaining addi-20 tional information concerning the deter-21 mination: and 22 "(iii) notification of the right to ap-23 peal the determination and instructions on 24 how to initiate an appeal in accordance 25 with subsection (d).

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"(c) GRIEVANCES.—A group health plan or a health
 insurance issuer shall have written procedures for address ing grievances between the plan or issuer offering health
 insurance coverage in connection with a group health plan
 and a participant or beneficiary. Determinations under
 such procedures shall be non-appealable.

7 "(d) INTERNAL APPEAL OF COVERAGE DETERMINA8 TIONS.—

9 "(1) RIGHT TO APPEAL.—

10 "(A) IN GENERAL.—A participant or bene-11 ficiary (or the authorized representative of the 12 participant or beneficiary) or the treating health 13 care professional with the consent of the partici-14 pant or beneficiary (or the authorized represent-15 ative of the participant or beneficiary), may ap-16 peal any adverse coverage determination under 17 subsection (b) under the procedures described in 18 this subsection.

19 "(B) TIME FOR APPEAL.—A plan or issuer
20 shall ensure that a participant or beneficiary
21 has a period of not less than 180 days beginning
22 on the date of an adverse coverage determination
23 under subsection (b) in which to appeal such de24 termination under this subsection.

1	"(C) FAILURE TO ACT.—The failure of a
2	plan or issuer to issue a determination under
3	subsection (b) within the applicable timeline es-
4	tablished for such a determination under such
5	subsection shall be treated as an adverse coverage
6	determination for purposes of proceeding to in-
7	ternal review under this subsection.
8	"(2) Records.—A group health plan and a
9	health insurance issuer shall maintain written
10	records, for at least 6 years, with respect to any ap-
11	peal under this subsection for purposes of internal
12	quality assurance and improvement. Nothing in the
13	preceding sentence shall be construed as preventing a
14	plan and issuer from entering into an agreement
15	under which the issuer agrees to assume responsibility
16	for compliance with the requirements of this section
17	and the plan is released from liability for such com-
18	pliance.

19 "(3) ROUTINE DETERMINATIONS.—A group
20 health plan or a health insurance issuer shall com21 plete the consideration of an appeal of an adverse
22 routine determination under this subsection not later
23 than 30 working days after the date on which a re24 quest for such appeal is received.

25 "(4) Expedited determination.—

1	"(A) IN GENERAL.—An expedited deter-
2	mination with respect to an appeal under this
3	subsection shall be made in accordance with the
4	medical exigencies of the case, but in no case
5	more than 72 hours after the request for such ap-
6	peal is received by the plan or issuer under sub-
7	paragraph (B) or (C).
8	"(B) Request by participant or bene-
9	FICIARY.—A plan or issuer shall maintain pro-
10	cedures for expediting a prior authorization de-
11	termination under this subsection upon the re-
12	quest of a participant or beneficiary if, based on
13	such a request, the plan or issuer determines that
14	the normal time for making such a determina-
15	tion could seriously jeopardize the life or health
16	of the participant or beneficiary.
17	"(C) DOCUMENTATION BY HEALTH CARE
18	PROFESSIONAL.—A plan or issuer shall main-
19	tain procedures for expediting a prior authoriza-
20	tion determination under this subsection if the
21	request involved indicates that the treating
22	health care professional has reasonably docu-
23	mented, based on the medical exigencies of the
24	case that a determination under the procedures
25	described in paragraph (2) could seriously jeop-

ardize the life or health of the participant or 1 2 beneficiary. 3 "(5) CONDUCT OF REVIEW.—A review of an ad-4 verse coverage determination under this subsection 5 shall be conducted by an individual with appropriate 6 expertise who was not directly involved in the initial 7 determination. 8 "(6) LACK OF MEDICAL NECESSITY.—A review of 9 an appeal under this subsection relating to a deter-10 mination to deny coverage based on a lack of medical 11 necessity and appropriateness, or based on an experi-12 mental or investigational treatment, shall be made 13 only by a physician with appropriate expertise, in-14 cluding age-appropriate expertise, who was not involved in the initial determination. 15 "(7) NOTICE.— 16 17 "(A) IN GENERAL.—Written notice of a de-18 termination made under an internal review 19 process shall be issued to the participant or bene-20 ficiary (or the authorized representative of the 21 participant or beneficiary) and the treating 22 health care professional not later than 2 working

24 in the 72-hour period referred to in paragraph
25 (4) if applicable).

days after the completion of the review (or with-

1	"(B) Adverse coverage determina-
2	TIONS.—With respect to an adverse coverage de-
3	termination made under this subsection, the no-
4	tice described in subparagraph (A) shall
5	include—
6	"(i) the reasons for the determination
7	(including the clinical or scientific-evidence
8	based rationale used in making the deter-
9	mination) written in a manner to be under-
10	standable to the average participant or ben-
11	eficiary;
12	"(ii) the procedures for obtaining addi-
13	tional information concerning the deter-
14	mination; and
15	"(iii) notification of the right to an
16	independent external review under sub-
17	section (e) and instructions on how to ini-
18	tiate such a review.
19	"(e) INDEPENDENT EXTERNAL REVIEW.—
20	"(1) Access to review.—
21	"(A) IN GENERAL.—A group health plan or
22	a health insurance issuer offering health insur-
23	ance coverage in connection with a group health
24	plan shall have written procedures to permit a
25	participant or beneficiary (or the authorized rep-

1	resentative of the participant or beneficiary) ac-
2	cess to an independent external review with re-
3	spect to an adverse coverage determination con-
4	cerning a particular item or service (including a
5	circumstance treated as an adverse coverage de-
6	termination under subparagraph (B)) where—
7	"(i) the particular item or service
8	involved—
9	((I)(aa) would be a covered ben-
10	efit, when medically necessary and ap-
11	propriate under the terms and condi-
12	tions of the plan, and the item or serv-
13	ice has been determined not to be medi-
14	cally necessary and appropriate under
15	the internal appeals process required
16	under subsection (d) or there has been
17	a failure to issue a coverage determina-
18	tion as described in subparagraph (B) ;
19	and
20	"(bb)(AA) the amount of such
21	item or service involved exceeds a sig-
22	nificant financial threshold; or
23	"(BB) there is a significant risk
24	of placing the life or health of the par-
25	ticipant or beneficiary in jeopardy; or

1	"(II) would be a covered benefit,
2	when not considered experimental or
3	investigational under the terms and
4	conditions of the plan, and the item or
5	service has been determined to be ex-
6	perimental or investigational under the
7	internal appeals process required
8	under subsection (d) or there has been
9	a failure to issue a coverage determina-
10	tion as described in subparagraph (B) ;
11	and
12	"(ii) the participant or beneficiary has
13	completed the internal appeals process
14	under subsection (d) with respect to such de-
15	termination.
16	"(B) FAILURE TO ACT.—The failure of a
17	plan or issuer to issue a coverage determination
18	under subsection $(d)(6)$ within the applicable
19	timeline established for such a determination
20	under such subsection shall be treated as an ad-
21	verse coverage determination for purposes of pro-
22	ceeding to independent external review under
23	this subsection.
24	"(2) INITIATION OF THE INDEPENDENT EXTER-
25	NAL REVIEW PROCESS.—

1	"(A) FILING OF REQUEST.—A participant
2	or beneficiary (or the authorized representative
3	of the participant or beneficiary) who desires to
4	have an independent external review conducted
5	under this subsection shall file a written request
6	for such a review with the plan or issuer in-
7	volved not later than 30 working days after the
8	receipt of a final denial of a claim under sub-
9	section (d). Any such request shall include the
10	consent of the participant or beneficiary (or the
11	authorized representative of the participant or
12	beneficiary) for the release of medical informa-
13	tion and records to independent external review-
14	ers regarding the participant or beneficiary.
15	"(B) TIMEFRAME FOR SELECTION OF AP-
16	PEALS ENTITY.—Not later than 5 working days
17	after the receipt of a request under subparagraph
18	(A), or earlier in accordance with the medical
19	exigencies of the case, the plan or issuer involved
20	shall—
21	"(i) select an external appeals entity
22	under paragraph $(3)(A)$ that shall be re-
23	sponsible for designating an independent ex-
24	ternal reviewer under paragraph $(3)(B);$
25	and

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1	"(ii) provide notice of such selection to
2	the participant or beneficiary (which shall
3	include the name and address of the entity).
4	"(C) Provision of information.—Not
5	later than 5 working days after the plan or
6	issuer provides the notice required under sub-
7	paragraph (B)(ii), or earlier in accordance with
8	the medical exigencies of the case, the plan,
9	issuer, participant, beneficiary or physician (of
10	the participant or beneficiary) involved shall for-
11	ward necessary information (including, only in
12	the case of a plan or issuer, medical records, any
13	relevant review criteria, the clinical rationale
14	consistent with the terms and conditions of the
15	contract between the plan or issuer and the par-
16	ticipant or beneficiary for the coverage denial,
17	and evidence of the coverage of the participant or
18	beneficiary) to the qualified external appeals en-
19	tity designated under paragraph $(3)(A)$.
20	"(D) Follow-up written notifica-
21	TION.—The plan or issuer involved shall send a
22	follow-up written notification, in a timely man-
23	ner, to the participant or beneficiary (or the au-
24	thorized representative of the participant or ben-
25	eficiary) and the plan administrator, indicating

1	that an independent external review has been
2	initiated.
3	"(3) Conduct of independent external re-
4	VIEW.—
5	"(A) DESIGNATION OF EXTERNAL APPEALS
6	ENTITY BY PLAN OR ISSUER.—
7	"(i) In general.—A plan or issuer
8	that receives a request for an independent
9	external $review$ $under$ $paragraph$ (2)(A)
10	shall designate a qualified entity described
11	in clause (ii), in a manner designed to en-
12	sure that the entity so designated will make
13	a decision in an unbiased manner, to serve
14	as the external appeals entity.
15	"(ii) Qualified entities.—A quali-
16	fied entity shall be—
17	"(I) an independent external re-
18	view entity licensed or credentialed by
19	a State;
20	"(II) a State agency established
21	for the purpose of conducting inde-
22	pendent external reviews;
23	"(III) any entity under contract
24	with the Federal Government to pro-

1	vide independent external review serv-
2	ices;
3	"(IV) any entity accredited as an
4	independent external review entity by
5	an accrediting body recognized by the
6	Secretary for such purpose; or
7	"(V) any other entity meeting cri-
8	teria established by the Secretary for
9	purposes of this subparagraph.
10	"(B) DESIGNATION OF INDEPENDENT EX-
11	TERNAL REVIEWER BY EXTERNAL APPEALS ENTI-
12	TY.—The external appeals entity designated
13	under subparagraph (A) shall, not later than 30
14	days after the date on which such entity is des-
15	ignated under subparagraph (A), or earlier in
16	accordance with the medical exigencies of the
17	case, designate one or more individuals to serve
18	as independent external reviewers with respect to
19	a request received under paragraph $(2)(A)$. Such
20	reviewers shall be independent medical experts
21	who shall—
22	((i) be appropriately credentialed or
23	licensed in any State to deliver health care
24	services;

1	"(ii) not have any material, profes-
2	sional, familial, or financial affiliation
3	with the case under review, the participant
4	or beneficiary involved, the treating health
5	care professional, the institution where the
6	treatment would take place, or the manufac-
7	turer of any drug, device, procedure, or
8	other therapy proposed for the participant
9	or beneficiary whose treatment is under re-
10	view;
11	"(iii) have expertise (including age-ap-
12	propriate expertise) in the diagnosis or
13	treatment under review and be a physician
14	of the same specialty, when reasonably
15	available, as the physician treating the par-
16	ticipant or beneficiary or recommending or
17	prescribing the treatment in question;
18	"(iv) receive only reasonable and cus-
19	tomary compensation from the group health
20	plan or health insurance issuer in connec-
21	tion with the independent external review
22	that is not contingent on the decision ren-
23	dered by the reviewer; and
24	"(v) not be held liable for decisions re-
25	garding medical determinations (but may
1	be held liable for actions that are arbitrary
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2	and capricious).
3	"(4) Standard of review.—
4	"(A) IN GENERAL.—An independent exter-
5	nal reviewer shall—
6	"(i) make an independent determina-
7	tion based on the valid, relevant, scientific
8	and clinical evidence to determine the med-
9	ical necessity, appropriateness, experi-
10	mental or investigational nature of the pro-
11	posed treatment; and
12	"(ii) take into consideration appro-
13	priate and available information, including
14	any evidence-based decision making or clin-
15	ical practice guidelines used by the group
16	health plan or health insurance issuer;
17	timely evidence or information submitted by
18	the plan, issuer, patient or patient's physi-
19	cian; the patient's medical record; expert
20	consensus including both generally accepted
21	medical practice and recognized best prac-
22	tice; medical literature as defined in section
23	556(5) of the Federal Food, Drug, and Cos-
24	metic Act; the following standard reference
25	compendia: The American Hospital For-

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1	mulary Service-Drug Information, the
2	American Dental Association Accepted Den-
3	tal Therapeutics, and the United States
4	Pharmacopoeia-Drug Information; and
5	findings, studies, or research conducted by
6	or under the auspices of Federal Govern-
7	ment agencies and nationally recognized
8	Federal research institutes including the
9	Agency for Healthcare Research and Qual-
10	ity, National Institutes of Health, National
11	Academy of Sciences, Health Care Financ-
12	ing Administration, and any national
13	board recognized by the National Institutes
14	of Health for the purposes of evaluating the
15	medical value of health services.
16	"(B) NOTICE.—The plan or issuer involved
17	shall ensure that the participant or beneficiary
18	receives notice, within 30 days after the deter-
19	mination of the independent medical expert, re-
20	garding the actions of the plan or issuer with re-
21	spect to the determination of such expert under
22	the independent external review.
23	"(5) TIMEFRAME FOR REVIEW.—
24	"(A) IN GENERAL.—The independent exter-
25	nal reviewer shall complete a review of an ad-

1	verse coverage determination in accordance with
2	the medical exigencies of the case.
3	"(B) Expedited review.—Notwith-
4	standing subparagraph (A), a review described
5	in such subparagraph shall be completed not
6	later than 72 hours after the later of—
7	"(i) the date on which such reviewer is
8	designated; or
9	"(ii) the date on which all information
10	necessary to completing such review is re-
11	ceived;
12	if the completion of such review in a period of
13	time in excess of 72 hours would seriously jeop-
14	ardize the life or health of the participant or
15	beneficiary.
16	"(C) LIMITATION.—Notwithstanding sub-
17	paragraph (A), and except as provided in sub-
18	paragraph (B), a review described in subpara-
19	graph (A) shall be completed not later than 30
20	working days after the later of—
21	"(i) the date on which such reviewer is
22	designated; or
23	"(ii) the date on which all information
24	necessary to completing such review is re-
25	ceived.

1	"(6) Binding determination and access to
2	CARE.—
3	"(A) IN GENERAL.—The determination of
4	an independent external reviewer under this sub-
5	section shall be binding upon the plan or issuer
6	if the provisions of this subsection or the proce-
7	dures implemented under such provisions were
8	complied with by the independent external re-
9	viewer.
10	$('/\mathbf{D})$ Therefore for comparison of

(B) TIMETABLE FOR COMMENCEMENT OF 10 11 CARE.—Where an independent external reviewer 12 determines that the participant or beneficiary is 13 entitled to coverage of the items or services that 14 were the subject of the review, the reviewer shall 15 establish a timeframe, in accordance with the 16 medical exigencies of the case, during which the 17 plan or issuer shall comply with the decision of 18 the reviewer with respect to the coverage of such 19 items or services under the terms and conditions 20 of the plan.

21 "(C) FAILURE TO COMPLY.—If a plan or
22 issuer fails to comply with the timeframe estab23 lished under subparagraph (B) with respect to a
24 participant or beneficiary, where such failure to
25 comply is caused by the plan or issuer, the par-

1	ticipant or beneficiary may obtain the items or
2	services involved (in a manner consistent with
3	the determination of the independent external re-
4	viewer) from any provider regardless of whether
5	such provider is a participating provider under
6	the plan or coverage.
7	"(D) Reimbursement.—
8	"(i) IN GENERAL.—Where a partici-
9	pant or beneficiary obtains items or services
10	in accordance with subparagraph (C), the
11	plan or issuer involved shall provide for re-
12	imbursement of the costs of such items of
13	services. Such reimbursement shall be made
14	to the treating provider or to the partici-
15	pant or beneficiary (in the case of a partici-
16	pant or beneficiary who pays for the costs
17	of such items or services).
18	"(ii) Amount.—The plan or issuer
19	shall fully reimburse a provider, partici-
20	pant or beneficiary under clause (i) for the
21	total costs of the items or services provided
22	(regardless of any plan limitations that
23	may apply to the coverage of such items of
24	services) so long as—

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1	((I) the items or services would
2	have been covered under the terms of
3	the plan or coverage if provided by the
4	plan or issuer; and
5	"(II) the items or services were
6	provided in a manner consistent with
7	the determination of the independent
8	external reviewer.
9	"(E) FAILURE TO REIMBURSE.—Where a
10	plan or issuer fails to provide reimbursement to
11	a provider, participant or beneficiary in accord-
12	ance with this paragraph, the provider, partici-
13	pant or beneficiary may commence a civil action
14	(or utilize other remedies available under law) to
15	recover only the amount of any such reimburse-
16	ment that is unpaid and any necessary legal
17	costs or expenses (including attorneys' fees) in-
18	curred in recovering such reimbursement.
19	"(7) STUDY.—Not later than 2 years after the
20	date of enactment of this section, the General Ac-
21	counting Office shall conduct a study of a statistically
22	appropriate sample of completed independent external
23	reviews. Such study shall include an assessment of the
24	process involved during an independent external re-
25	view and the basis of decisionmaking by the inde-

pendent external reviewer. The results of such study
 shall be submitted to the appropriate committees of
 Congress.

4 "(8) EFFECT ON CERTAIN PROVISIONS.—Nothing
5 in this section shall be construed as affecting or modi6 fying section 514 of this Act with respect to a group
7 health plan.

8 "(f) RULE OF CONSTRUCTION.—Nothing in this sec-9 tion shall be construed to prohibit a plan administrator or 10 plan fiduciary or health plan medical director from request-11 ing an independent external review by an independent ex-12 ternal reviewer without first completing the internal review 13 process.

14 "(g) DEFINITIONS.—In this section:

15 "(1) ADVERSE COVERAGE DETERMINATION.—The
16 term 'adverse coverage determination' means a cov17 erage determination under the plan which results in
18 a denial of coverage or reimbursement.

19 "(2) COVERAGE DETERMINATION.—The term
20 'coverage determination' means with respect to items
21 and services for which coverage may be provided
22 under a health plan, a determination of whether or
23 not such items and services are covered or reimburs24 able under the coverage and terms of the contract.

"(3) GRIEVANCE.—The term 'grievance' means
any complaint made by a participant or beneficiary
that does not involve a coverage determination.
"(4) GROUP HEALTH PLAN.—The term 'group
health plan' shall have the meaning given such term
in section 733(a). In applying this paragraph, ex-
cepted benefits described in section 733(c) shall not be
treated as benefits consisting of medical care.
"(5) Health insurance coverage.—The term
'health insurance coverage' has the meaning given
such term in section 733(b)(1). In applying this
paragraph, excepted benefits described in section
733(c) shall not be treated as benefits consisting of

medical care.

15 "(6) HEALTH INSURANCE ISSUER.—The term
16 health insurance issuer' has the meaning given such
17 term in section 733(b)(2).

18 "(7) PRIOR AUTHORIZATION DETERMINATION.—
19 The term 'prior authorization determination' means a
20 coverage determination prior to the provision of the
21 items and services as a condition of coverage of the
22 items and services under the coverage.

23 "(8) TREATING HEALTH CARE PROFESSIONAL.—
24 The term 'treating health care professional' with re25 spect to a group health plan, health insurance issuer

or provider sponsored organization means a physi cian (medical doctor or doctor of osteopathy) or other
 health care practitioner who is acting within the
 scope of his or her State licensure or certification for
 the delivery of health care services and who is pri marily responsible for delivering those services to the
 participant or beneficiary.

8 "(9) UTILIZATION REVIEW.—The term 'utiliza-9 tion review' with respect to a group health plan or 10 health insurance coverage means a set of formal tech-11 niques designed to monitor the use of, or evaluate the 12 clinical necessity, appropriateness, efficacy, or effi-13 ciency of, health care services, procedures, or settings. 14 Techniques may include ambulatory review, prospec-15 tive review, second opinion, certification, concurrent 16 review, case management, discharge planning or ret-17 rospective review.".

18 (b) ENFORCEMENT.—Section 502(c) of the Employee 19 Retirement Income Security Act of 1974 (29 U.S.C. 1132(c)) is amended by adding at the end the following: 20 21 "(8) The Secretary may assess a civil penalty against 22 any plan of up to \$10,000 for the plan's failure or refusal 23 to comply with any timeline applicable under section 24 503(e) or any determination under such section, except that in any case in which treatment was not commenced by the 25

1 plan in accordance with the determination of an independent external reviewer, the Secretary shall assess a civil 2 penalty of \$10,000 against the plan and the plan shall pay 3 4 such penalty to the participant or beneficiary involved.". (c) CONFORMING AMENDMENT.—The table of contents 5 in section 1 of the Employee Retirement Income Security 6 7 Act of 1974 is amended by striking the item relating to 8 section 503 and inserting the following new item: "Sec. 503. Claims procedures, coverage determination, grievances and appeals.".

9 (d) EFFECTIVE DATE.—The amendments made by this 10 section shall apply with respect to plan years beginning on 11 or after 1 year after the date of enactment of this Act. The 12 Secretary shall issue all regulations necessary to carry out 13 the amendments made by this section before the effective 14 date thereof.

15 *TITLE II—WOMEN'S HEALTH AND* 16 *CANCER RIGHTS*

17 SEC. 201. WOMEN'S HEALTH AND CANCER RIGHTS.

(a) SHORT TITLE.—This section may be cited as the
"Women's Health and Cancer Rights Act of 1999".

20 (b) FINDINGS.—Congress finds that—

21 (1) the offering and operation of health plans af-

22 fect commerce among the States;

23 (2) health care providers located in a State serve

- 24 patients who reside in the State and patients who re-
- 25 side in other States; and

1	(3) in order to provide for uniform treatment of
2	health care providers and patients among the States,
3	it is necessary to cover health plans operating in 1
4	State as well as health plans operating among the
5	several States.
6	(c) Amendments to ERISA.—
7	(1) IN GENERAL.—Subpart B of part 7 of sub-
8	title B of title I of the Employee Retirement Income
9	Security Act of 1974, as amended by section 111(a),
10	is further amended by adding at the end the fol-
11	lowing:
12	"SEC. 715. REQUIRED COVERAGE FOR MINIMUM HOSPITAL
13	STAY FOR MASTECTOMIES AND LYMPH NODE
13 14	STAY FOR MASTECTOMIES AND LYMPH NODE DISSECTIONS FOR THE TREATMENT OF
_	
14	DISSECTIONS FOR THE TREATMENT OF
14 15	DISSECTIONS FOR THE TREATMENT OF BREAST CANCER AND COVERAGE FOR SEC-
14 15 16	DISSECTIONS FOR THE TREATMENT OF BREAST CANCER AND COVERAGE FOR SEC- ONDARY CONSULTATIONS.
14 15 16 17	DISSECTIONS FOR THE TREATMENT OF BREAST CANCER AND COVERAGE FOR SEC- ONDARY CONSULTATIONS. "(a) INPATIENT CARE.—
14 15 16 17 18	DISSECTIONS FOR THE TREATMENT OF BREAST CANCER AND COVERAGE FOR SEC- ONDARY CONSULTATIONS. "(a) INPATIENT CARE.— "(1) IN GENERAL.—A group health plan, and a
14 15 16 17 18 19	DISSECTIONS FOR THE TREATMENT OF BREAST CANCER AND COVERAGE FOR SEC- ONDARY CONSULTATIONS. "(a) INPATIENT CARE.— "(1) IN GENERAL.—A group health plan, and a health insurance issuer providing health insurance
 14 15 16 17 18 19 20 	DISSECTIONS FOR THE TREATMENT OF BREAST CANCER AND COVERAGE FOR SEC- ONDARY CONSULTATIONS. "(a) INPATIENT CARE.— "(1) IN GENERAL.—A group health plan, and a health insurance issuer providing health insurance coverage in connection with a group health plan, that
14 15 16 17 18 19 20 21	DISSECTIONS FOR THE TREATMENT OF BREAST CANCER AND COVERAGE FOR SEC- ONDARY CONSULTATIONS. "(a) INPATIENT CARE.— "(1) IN GENERAL.—A group health plan, and a health insurance issuer providing health insurance coverage in connection with a group health plan, that provides medical and surgical benefits shall ensure

1	tion with the patient, to be medically necessary and
2	appropriate following—
3	"(A) a mastectomy;
4	"(B) a lumpectomy; or
5	``(C) a lymph node dissection for the treat-
6	ment of breast cancer.
7	"(2) EXCEPTION.—Nothing in this section shall
8	be construed as requiring the provision of inpatient
9	coverage if the attending physician and patient deter-
10	mine that a shorter period of hospital stay is medi-
11	cally appropriate.
12	"(b) Prohibition on Certain Modifications.—In
13	implementing the requirements of this section, a group
14	health plan, and a health insurance issuer providing health
15	insurance coverage in connection with a group health plan,
16	may not modify the terms and conditions of coverage based
17	on the determination by a participant or beneficiary to re-
18	quest less than the minimum coverage required under sub-
19	section (a).
20	"(c) NOTICE.—A group health plan, and a health in-
21	surance issuer providing health insurance coverage in con-
22	nection with a group health plan shall provide notice to

23 each participant and beneficiary under such plan regarding24 the coverage required by this section in accordance with reg-

1	be in writing and prominently positioned in any literature
2	or correspondence made available or distributed by the plan
3	or issuer and shall be transmitted—
4	"(1) in the next mailing made by the plan or
5	issuer to the participant or beneficiary;
6	"(2) as part of any yearly informational packet
7	sent to the participant or beneficiary; or
8	"(3) not later than January 1, 2000;
9	whichever is earlier.
10	"(d) Secondary Consultations.—
11	"(1) IN GENERAL.—A group health plan, and a
12	health insurance issuer providing health insurance
13	coverage in connection with a group health plan, that
14	provides coverage with respect to medical and sur-
15	gical services provided in relation to the diagnosis
16	and treatment of cancer shall ensure that full coverage
17	is provided for secondary consultations by specialists
18	in the appropriate medical fields (including pathol-
19	ogy, radiology, and oncology) to confirm or refute
20	such diagnosis. Such plan or issuer shall ensure that
21	full coverage is provided for such secondary consulta-
22	tion whether such consultation is based on a positive
23	or negative initial diagnosis. In any case in which
24	the attending physician certifies in writing that serv-
25	ices necessary for such a secondary consultation are

1	not sufficiently available from specialists operating
2	under the plan with respect to whose services coverage
3	is otherwise provided under such plan or by such
4	issuer, such plan or issuer shall ensure that coverage
5	is provided with respect to the services necessary for
6	the secondary consultation with any other specialist
7	selected by the attending physician for such purpose
8	at no additional cost to the individual beyond that
9	which the individual would have paid if the specialist
10	was participating in the network of the plan.
11	"(2) Exception.—Nothing in paragraph (1)
12	shall be construed as requiring the provision of sec-
13	ondary consultations where the patient determines not
14	to seek such a consultation.
15	"(e) Prohibition on Penalties or Incentives.—
16	A group health plan, and a health insurance issuer pro-
17	viding health insurance coverage in connection with a
18	group health plan, may not—
19	"(1) penalize or otherwise reduce or limit the re-
20	imbursement of a provider or specialist because the
21	provider or specialist provided care to a participant
22	or beneficiary in accordance with this section;
23	"(2) provide financial or other incentives to a
24	physician or specialist to induce the physician or spe-
25	cialist to keep the length of inpatient stays of patients

1	following a mastectomy, lumpectomy, or a lymph
2	node dissection for the treatment of breast cancer
3	below certain limits or to limit referrals for secondary
4	consultations; or
5	"(3) provide financial or other incentives to a

6 physician or specialist to induce the physician or spe7 cialist to refrain from referring a participant or bene8 ficiary for a secondary consultation that would other9 wise be covered by the plan or coverage involved
10 under subsection (d).".

11 (2) CLERICAL AMENDMENT.—The table of con12 tents in section 1 of the Employee Retirement Income
13 Security Act of 1974 is amended by inserting after
14 the item relating to section 714 the following new
15 item:

"Sec. 715. Required coverage for minimum hospital stay for mastectomies and lymph node dissections for the treatment of breast cancer and coverage for secondary consultations.".

16 (d) AMENDMENTS TO PHSA RELATING TO THE
17 GROUP MARKET.—Subpart 2 of part A of title XXVII of
18 the Public Health Service Act (42 U.S.C. 300gg-4 et seq.)
19 is amended by adding at the end the following new section:

1	"SEC. 2707. REQUIRED COVERAGE FOR MINIMUM HOSPITAL			
2	STAY FOR MASTECTOMIES AND LYMPH NODE			
3	DISSECTIONS FOR THE TREATMENT OF			
4	BREAST CANCER AND COVERAGE FOR SEC-			
5	ONDARY CONSULTATIONS.			
6	"(a) INPATIENT CARE.—			
7	"(1) IN GENERAL.—A group health plan, and a			
8	health insurance issuer providing health insurance			
9	coverage in connection with a group health plan, that			
10	provides medical and surgical benefits shall ensure			
11	that inpatient coverage with respect to the treatment			
12	of breast cancer is provided for a period of time as			
13	is determined by the attending physician, in consulta-			
14	tion with the patient, to be medically necessary and			
15	appropriate following—			
16	"(A) a mastectomy;			
17	"(B) a lumpectomy; or			
18	"(C) a lymph node dissection for the treat-			
19	ment of breast cancer.			
20	"(2) Exception.—Nothing in this section shall			
21	be construed as requiring the provision of inpatient			
22	coverage if the attending physician and patient deter-			
23	mine that a shorter period of hospital stay is medi-			
24	cally appropriate.			
25	"(b) Prohibition on Certain Modifications.—In			
26	implementing the requirements of this section, a group			

health plan, and a health insurance issuer providing health
 insurance coverage in connection with a group health plan,
 may not modify the terms and conditions of coverage based
 on the determination by a participant or beneficiary to re quest less than the minimum coverage required under sub section (a).

7 "(c) NOTICE.—A group health plan, and a health in-8 surance issuer providing health insurance coverage in con-9 nection with a group health plan shall provide notice to 10 each participant and beneficiary under such plan regarding 11 the coverage required by this section in accordance with regulations promulgated by the Secretary. Such notice shall 12 be in writing and prominently positioned in any literature 13 or correspondence made available or distributed by the plan 14 15 or issuer and shall be transmitted—

16 "(1) in the next mailing made by the plan or
17 issuer to the participant or beneficiary;

18 "(2) as part of any yearly informational packet

19 sent to the participant or beneficiary; or

20 "(3) not later than January 1, 2000;

21 whichever is earlier.

22 "(d) SECONDARY CONSULTATIONS.—

23 "(1) IN GENERAL.—A group health plan, and a
24 health insurance issuer providing health insurance
25 coverage in connection with a group health plan that

1	provides coverage with respect to medical and sur-
2	gical services provided in relation to the diagnosis
3	and treatment of cancer shall ensure that full coverage
4	is provided for secondary consultations by specialists
5	in the appropriate medical fields (including pathol-
6	ogy, radiology, and oncology) to confirm or refute
7	such diagnosis. Such plan or issuer shall ensure that
8	full coverage is provided for such secondary consulta-
9	tion whether such consultation is based on a positive
10	or negative initial diagnosis. In any case in which
11	the attending physician certifies in writing that serv-
12	ices necessary for such a secondary consultation are
13	not sufficiently available from specialists operating
14	under the plan with respect to whose services coverage
15	is otherwise provided under such plan or by such
16	issuer, such plan or issuer shall ensure that coverage
17	is provided with respect to the services necessary for
18	the secondary consultation with any other specialist
19	selected by the attending physician for such purpose
20	at no additional cost to the individual beyond that
21	which the individual would have paid if the specialist
22	was participating in the network of the plan.
23	"(2) EXCEPTION.—Nothing in paragraph (1)

- 24 shall be construed as requiring the provision of sec-
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ondary consultations where the patient determines not
 to seek such a consultation.

3 "(e) PROHIBITION ON PENALTIES OR INCENTIVES.—
4 A group health plan, and a health insurance issuer pro5 viding health insurance coverage in connection with a
6 group health plan, may not—

7 "(1) penalize or otherwise reduce or limit the re8 imbursement of a provider or specialist because the
9 provider or specialist provided care to a participant
10 or beneficiary in accordance with this section;

11 "(2) provide financial or other incentives to a 12 physician or specialist to induce the physician or spe-13 cialist to keep the length of inpatient stays of patients 14 following a mastectomy, lumpectomy, or a lymph 15 node dissection for the treatment of breast cancer 16 below certain limits or to limit referrals for secondary 17 consultations; or

18 "(3) provide financial or other incentives to a 19 physician or specialist to induce the physician or spe-20 cialist to refrain from referring a participant or bene-21 ficiary for a secondary consultation that would other-22 wise be covered by the plan or coverage involved 23 under subsection (d).".

24 (e) AMENDMENTS TO PHSA RELATING TO THE INDI25 VIDUAL MARKET.—The first subpart 3 of part B of title

XXVII of the Public Health Service Act (42 U.S.C. 300gg-1 2 51 et seq.) (relating to other requirements) (42 U.S.C. 3 300qq-51 et seq.) is amended— 4 (1) by redesignating such subpart as subpart 2; 5 and 6 (2) by adding at the end the following: 7 **"SEC. 2753. REQUIRED COVERAGE FOR MINIMUM HOSPITAL** 8 STAY FOR MASTECTOMIES AND LYMPH NODE 9 DISSECTIONS FOR THE TREATMENT OF 10 BREAST CANCER AND SECONDARY CON-11 SULTATIONS. 12 "The provisions of section 2707 shall apply to health 13 insurance coverage offered by a health insurance issuer in the individual market in the same manner as they apply 14 15 to health insurance coverage offered by a health insurance issuer in connection with a group health plan in the small 16 17 or large group market.".

18 (f) Amendments to the IRC.—

19 (1) IN GENERAL.—Subchapter B of chapter 100
20 of the Internal Revenue Code of 1986, as amended by
21 section 111(b), is further amended by inserting after
22 section 9813 the following:

1	"SEC. 9814. REQUIRED COVERAGE FOR MINIMUM HOSPITAL		
2	STAY FOR MASTECTOMIES AND LYMPH NODE		
3	DISSECTIONS FOR THE TREATMENT OF		
4	BREAST CANCER AND COVERAGE FOR SEC-		
5	ONDARY CONSULTATIONS.		
6	"(a) INPATIENT CARE.—		
7	"(1) In general.—A group health plan that		
8	B provides medical and surgical benefits shall ensur		
9	9 that inpatient coverage with respect to the treatmer		
10	0 of breast cancer is provided for a period of time a		
11	is determined by the attending physician, in consulta-		
12	tion with the patient, to be medically necessary and		
13	appropriate following—		
14	"(A) a mastectomy;		
15	"(B) a lumpectomy; or		
16	``(C) a lymph node dissection for the treat-		
17	ment of breast cancer.		
18	"(2) EXCEPTION.—Nothing in this section shall		
19	be construed as requiring the provision of inpatient		
20	coverage if the attending physician and patient deter-		
21	mine that a shorter period of hospital stay is medi-		
22	cally appropriate.		
23	"(b) Prohibition on Certain Modifications.—In		
24	implementing the requirements of this section, a group		
25	health plan may not modify the terms and conditions of		
26	coverage based on the determination by a participant or		
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beneficiary to request less than the minimum coverage re quired under subsection (a).

"(c) NOTICE.—A group health plan shall provide notice to each participant and beneficiary under such plan
regarding the coverage required by this section in accordance with regulations promulgated by the Secretary. Such
notice shall be in writing and prominently positioned in
any literature or correspondence made available or distributed by the plan and shall be transmitted—

"(1) in the next mailing made by the plan to the
participant or beneficiary;

12 "(2) as part of any yearly informational packet
13 sent to the participant or beneficiary; or

14 *"(3) not later than January 1, 2000;*

15 whichever is earlier.

16 "(d) SECONDARY CONSULTATIONS.—

17 "(1) IN GENERAL.—A group health plan that 18 provides coverage with respect to medical and sur-19 gical services provided in relation to the diagnosis 20 and treatment of cancer shall ensure that full coverage 21 is provided for secondary consultations by specialists 22 in the appropriate medical fields (including pathol-23 ogy, radiology, and oncology) to confirm or refute 24 such diagnosis. Such plan or issuer shall ensure that 25 full coverage is provided for such secondary consulta-

1	tion whether such consultation is based on a positive
2	or negative initial diagnosis. In any case in which
3	the attending physician certifies in writing that serv-
4	ices necessary for such a secondary consultation are
5	not sufficiently available from specialists operating
6	under the plan with respect to whose services coverage
7	is otherwise provided under such plan or by such
8	issuer, such plan or issuer shall ensure that coverage
9	is provided with respect to the services necessary for
10	the secondary consultation with any other specialist
11	selected by the attending physician for such purpose
12	at no additional cost to the individual beyond that
13	which the individual would have paid if the specialist
14	was participating in the network of the plan.
15	"(2) Exception.—Nothing in paragraph (1)
16	shall be construed as requiring the provision of sec-
17	ondary consultations where the patient determines not
18	to seek such a consultation.
19	"(e) PROHIBITION ON PENALTIES.—A group health

19 "(e) PROHIBITION ON PENALTIES.—A group health
20 plan may not—

21 "(1) penalize or otherwise reduce or limit the re22 imbursement of a provider or specialist because the
23 provider or specialist provided care to a participant
24 or beneficiary in accordance with this section;

1	"(2) provide financial or other incentives to a
2	physician or specialist to induce the physician or spe-
3	cialist to keep the length of inpatient stays of patients
4	following a mastectomy, lumpectomy, or a lymph
5	node dissection for the treatment of breast cancer
6	below certain limits or to limit referrals for secondary
7	consultations; or
8	"(3) provide financial or other incentives to a
9	physician or specialist to induce the physician or spe-
10	cialist to refrain from referring a participant or bene-
11	ficiary for a secondary consultation that would other-
12	wise be covered by the plan involved under subsection
13	<i>(d)."</i> .
14	(2) Clerical Amendment.—The table of con-
15	tents for chapter 100 of such Code is amended by in-
16	serting after the item relating to section 9813 the fol-
17	lowing new item:
	"Sec. 9814. Required coverage for minimum hospital stay for mastectomies and lymph node dissections for the treatment of breast cancer and coverage for secondary consultations.".
18	TITLE III—GENETIC
19	INFORMATION AND SERVICES
20	SEC. 301. SHORT TITLE.
21	This title may be cited as the "Genetic Information

22 Nondiscrimination in Health Insurance Act of 1999".

1SEC. 302. AMENDMENTS TO EMPLOYEE RETIREMENT IN-2COME SECURITY ACT OF 1974.

3 (a) PROHIBITION OF HEALTH DISCRIMINATION ON
4 THE BASIS OF GENETIC INFORMATION OR GENETIC SERV5 ICES.—

6 (1) NO ENROLLMENT RESTRICTION FOR GENETIC
7 SERVICES.—Section 702(a)(1)(F) of the Employee Re8 tirement Income Security Act of 1974 (29 U.S.C.
9 1182(a)(1)(F)) is amended by inserting before the pe10 riod the following: "(including information about a
11 request for or receipt of genetic services)".

(2) NO DISCRIMINATION IN GROUP PREMIUMS
BASED ON PREDICTIVE GENETIC INFORMATION.—Subpart B of part 7 of subtitle B of title I of the Employee Retirement Income Security Act of 1974, as
amended by sections 111(a) and 201, is further
amended by adding at the end the following:

18 "SEC.716.PROHIBITINGPREMIUMDISCRIMINATION19AGAINST GROUPS ON THE BASIS OF PRE-20DICTIVE GENETIC INFORMATION.

21 "A group health plan, or a health insurance issuer of22 fering group health insurance coverage in connection with
23 a group health plan, shall not adjust premium or contribu24 tion amounts for a group on the basis of predictive genetic
25 information concerning any individual (including a de26 pendent) or family member of the individual (including inHR 2990 EAS

formation about a request for or receipt of genetic serv ices).".

- 3 (3) Conforming Amendments.— 4 (A) IN GENERAL.—Section 702(b) of the 5 Employee Retirement Income Security Act of 6 1974 (29 U.S.C. 1182(b)) is amended by adding 7 at the end the following: 8 "(3) Reference to related provision.—For 9 a provision prohibiting the adjustment of premium or 10 contribution amounts for a group under a group 11 health plan on the basis of predictive genetic informa-12 tion (including information about a request for or re-13 ceipt of genetic services), see section 716.". 14 (B) TABLE OF CONTENTS.—The table of 15 contents in section 1 of the Employee Retirement 16 Income Security Act of 1974, as amended by sec-17 tions 111(a) and 201, is further amended by in-18 serting after the item relating to section 715 the
- 19 following new item:

"Sec. 716. Prohibiting premium discrimination against groups on the basis of predictive genetic information.".

(b) LIMITATION ON COLLECTION OF PREDICTIVE GE21 NETIC INFORMATION.—Section 702 of the Employee Retire22 ment Income Security Act of 1974 (29 U.S.C. 1182) is
23 amended by adding at the end the following:

"(c) Collection of Predictive Genetic Informa Tion.—

3	"(1) Limitation on requesting or requiring
4	PREDICTIVE GENETIC INFORMATION.—Except as pro-
5	vided in paragraph (2), a group health plan, or a
6	health insurance issuer offering health insurance cov-
7	erage in connection with a group health plan, shall
8	not request or require predictive genetic information
9	concerning any individual (including a dependent) or
10	family member of the individual (including informa-
11	tion about a request for or receipt of genetic services).
12	"(2) INFORMATION NEEDED FOR DIAGNOSIS,
13	TREATMENT, OR PAYMENT.—
14	"(A) IN GENERAL.—Notwithstanding para-
15	graph (1), a group health plan, or a health in-
16	surance issuer offering health insurance coverage
17	in connection with a group health plan, that
18	provides health care items and services to an in-
19	dividual or dependent may request (but may not
20	require) that such individual or dependent dis-
21	close, or authorize the collection or disclosure of,
22	predictive genetic information for purposes of di-
23	agnosis, treatment, or payment relating to the
24	provision of health care items and services to
25	such individual or dependent.

"(B) NOTICE OF CONFIDENTIALITY PRAC-1 2 TICES AND DESCRIPTION OF SAFEGUARDS.—As a 3 part of a request under subparagraph (A), the 4 group health plan, or a health insurance issuer 5 offering health insurance coverage in connection 6 with a group health plan, shall provide to the in-7 dividual or dependent a description of the proce-8 dures in place to safeguard the confidentiality, 9 as described in subsection (d), of such predictive 10 genetic information. 11 "(d) Confidentiality with Respect to Pre-DICTIVE GENETIC INFORMATION.— 12 13 "(1) Notice of confidentiality practices.— 14 "(A) PREPARATION OF WRITTEN NOTICE.— 15 A group health plan, or a health insurance 16 issuer offering health insurance coverage in con-17 nection with a group health plan, shall post or 18 provide, in writing and in a clear and con-19 spicuous manner, notice of the plan or issuer's 20 confidentiality practices, that shall include— 21 "(i) a description of an individual's 22 rights with respect to predictive genetic in-23 formation;

1	"(ii) the procedures established by the
2	plan or issuer for the exercise of the individ-
3	ual's rights; and
4	"(iii) the right to obtain a copy of the
5	notice of the confidentiality practices re-
6	quired under this subsection.
7	"(B) MODEL NOTICE.—The Secretary, in
8	consultation with the National Committee on
9	Vital and Health Statistics and the National As-
10	sociation of Insurance Commissioners, and after
11	notice and opportunity for public comment, shall
12	develop and disseminate model notices of con-
13	fidentiality practices. Use of the model notice
14	shall serve as a defense against claims of receiv-
15	ing inappropriate notice.
16	"(2) ESTABLISHMENT OF SAFEGUARDS.—A
17	group health plan, or a health insurance issuer offer-
18	ing health insurance coverage in connection with a
19	group health plan, shall establish and maintain ap-
20	propriate administrative, technical, and physical
21	safeguards to protect the confidentiality, security, ac-
22	curacy, and integrity of predictive genetic informa-
23	tion created, received, obtained, maintained, used,
24	transmitted, or disposed of by such plan or issuer.".

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1	(c) DEFINITIONS.—Section 733(d) of the Employee Re-
2	tirement Income Security Act of 1974 (29 U.S.C. 1191b(d))
3	is amended by adding at the end the following:
4	"(5) FAMILY MEMBER.—The term 'family mem-
5	ber' means with respect to an individual—
6	"(A) the spouse of the individual;
7	"(B) a dependent child of the individual,
8	including a child who is born to or placed for
9	adoption with the individual; and
10	(C) all other individuals related by blood
11	to the individual or the spouse or child described
12	in subparagraph (A) or (B).
13	"(6) GENETIC INFORMATION.—The term 'genetic
14	information' means information about genes, gene
15	products, or inherited characteristics that may derive
16	from an individual or a family member (including
17	information about a request for or receipt of genetic
18	services).
19	"(7) GENETIC SERVICES.—The term 'genetic
20	services' means health services provided to obtain, as-
21	sess, or interpret genetic information for diagnostic
22	and therapeutic purposes, and for genetic education
23	and counseling.
24	"(8) Predictive genetic information.—

1	"(A) IN GENERAL.—The term 'predictive ge-
2	netic information' means, in the absence of
3	symptoms, clinical signs, or a diagnosis of the
4	condition related to such information—
5	"(i) information about an individual's
6	genetic tests;
7	"(ii) information about genetic tests of
8	family members of the individual; or
9	"(iii) information about the occurrence
10	of a disease or disorder in family members.
11	"(B) EXCEPTIONS.—The term 'predictive
12	genetic information' shall not include—
13	"(i) information about the sex or age of
14	the individual;
15	"(ii) information derived from phys-
16	ical tests, such as the chemical, blood, or
17	urine analyses of the individual including
18	cholesterol tests; and
19	"(iii) information about physical
20	exams of the individual.
21	"(9) GENETIC TEST.—The term 'genetic test'
22	means the analysis of human DNA, RNA, chro-
23	mosomes, proteins, and certain metabolites, including
24	analysis of genotypes, mutations, phenotypes, or
25	karyotypes, for the purpose of predicting risk of dis-

1 ease in asymptomatic or undiagnosed individuals. 2 Such term does not include physical tests, such as the 3 chemical, blood, or urine analyses of the individual 4 including cholesterol tests, and physical exams of the individual, in order to detect symptoms, clinical 5 6 signs, or a diagnosis of disease.". 7 (d) EFFECTIVE DATE.—Except as provided in this sec-8 tion, this section and the amendments made by this section 9 shall apply with respect to group health plans for plan 10 years beginning 1 year after the date of the enactment of 11 this Act. SEC. 303. AMENDMENTS TO THE PUBLIC HEALTH SERVICE 12 13 ACT. 14 (a) Amendments Relating to the Group Mar-15 KET.— 16 (1) **Prohibition of health discrimination** 17 ON THE BASIS OF GENETIC INFORMATION IN THE 18 GROUP MARKET.— 19 (A) NO ENROLLMENT RESTRICTION FOR GE-20 NETIC SERVICES.—Section 2702(a)(1)(F) of the 21 Public Health Service Act (42 U.S.C. 300gg-22 1(a)(1)(F) is amended by inserting before the 23 period the following: "(including information 24 about a request for or receipt of genetic serv-25 ices)".

1		(B) NO DISCRIMINATION IN PREMIUMS
2		BASED ON PREDICTIVE GENETIC INFORMATION.—
3		Subpart 2 of part A of title XXVII of the Public
4		Health Service Act, as amended by section 201,
5		is further amended by adding at the end the fol-
6		lowing new section:
7	"SEC.	2708. PROHIBITING PREMIUM DISCRIMINATION
8		AGAINST GROUPS ON THE BASIS OF PRE-

9 DICTIVE GENETIC INFORMATION IN THE 10 GROUP MARKET.

11 "A group health plan, or a health insurance issuer of-12 fering group health insurance coverage in connection with 13 a group health plan shall not adjust premium or contribu-14 tion amounts for a group on the basis of predictive genetic 15 information concerning any individual (including a de-16 pendent) or family member of the individual (including in-17 formation about a request for or receipt of genetic serv-18 ices).".

19	(C) Conforming Amendment.—Section
20	2702(b) of the Public Health Service Act (42
21	U.S.C. 300gg–1(b)) is amended by adding at the
22	end the following:
23	"(3) Reference to related provision.—For

a provision prohibiting the adjustment of premium orcontribution amounts for a group under a group

1	health plan on the basis of predictive genetic informa-
2	tion (including information about a request for or re-
3	ceipt of genetic services), see section 2708.".
4	(D) LIMITATION ON COLLECTION AND DIS-
5	CLOSURE OF PREDICTIVE GENETIC INFORMA-
6	TION.—Section 2702 of the Public Health Service
7	Act (42 U.S.C. $300gg-1$) is amended by adding
8	at the end the following:
9	"(c) Collection of Predictive Genetic Informa-
10	TION.—
11	"(1) Limitation on requesting or requiring
12	PREDICTIVE GENETIC INFORMATION.—Except as pro-
13	vided in paragraph (2), a group health plan, or a
14	health insurance issuer offering health insurance cov-
15	erage in connection with a group health plan, shall
16	not request or require predictive genetic information
17	concerning any individual (including a dependent) or
18	a family member of the individual (including infor-
19	mation about a request for or receipt of genetic serv-
20	ices).
21	"(2) Information needed for diagnosis,
22	TREATMENT, OR PAYMENT.—
23	"(A) IN GENERAL.—Notwithstanding para-
24	graph (1), a group health plan, or a health in-
25	surance issuer offering health insurance coverage

1	in connection with a group health plan, that
2	provides health care items and services to an in-
3	dividual or dependent may request (but may not
4	require) that such individual or dependent dis-
5	close, or authorize the collection or disclosure of,
6	predictive genetic information for purposes of di-
7	agnosis, treatment, or payment relating to the
8	provision of health care items and services to
9	such individual or dependent.
10	"(B) Notice of confidentiality prac-
11	tices and description of safeguards.—As a
12	part of a request under subparagraph (A) , the
13	group health plan, or a health insurance issuer
14	offering health insurance coverage in connection
15	with a group health plan, shall provide to the in-
16	dividual or dependent a description of the proce-
17	dures in place to safeguard the confidentiality,
18	as described in subsection (d), of such predictive
19	genetic information.
20	"(d) Confidentiality with Respect to Pre-
21	DICTIVE GENETIC INFORMATION.—
22	"(1) Notice of confidentiality practices.—
23	"(A) Preparation of written notice.—
24	A group health plan, or a health insurance
25	issuer offering health insurance coverage in con-

1	nection with a group health plan, shall post or
2	provide, in writing and in a clear and con-
3	spicuous manner, notice of the plan or issuer's
4	confidentiality practices, that shall include—
5	"(i) a description of an individual's
6	rights with respect to predictive genetic in-
7	formation;
8	``(ii) the procedures established by the
9	plan or issuer for the exercise of the individ-
10	ual's rights; and
11	"(iii) the right to obtain a copy of the
12	notice of the confidentiality practices re-
13	quired under this subsection.
14	"(B) MODEL NOTICE.—The Secretary, in
15	consultation with the National Committee on
16	Vital and Health Statistics and the National As-
17	sociation of Insurance Commissioners, and after
18	notice and opportunity for public comment, shall
19	develop and disseminate model notices of con-
20	fidentiality practices. Use of the model notice
21	shall serve as a defense against claims of receiv-
22	ing inappropriate notice.
23	"(2) ESTABLISHMENT OF SAFEGUARDS.—A
24	group health plan, or a health insurance issuer offer-
25	ing health insurance coverage in connection with a
group health plan, shall establish and maintain ap-	
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propriate administrative, technical, and physical	
safeguards to protect the confidentiality, security, ac-	
curacy, and integrity of predictive genetic informa-	
tion created, received, obtained, maintained, used,	
transmitted, or disposed of by such plan or issuer.".	
(2) DEFINITIONS.—Section 2791(d) of the Public	
Health Service Act (42 U.S.C. $300gg-91(d)$) is	
amended by adding at the end the following:	
"(15) FAMILY MEMBER.—The term 'family mem-	
ber' means, with respect to an individual—	
"(A) the spouse of the individual;	
``(B) a dependent child of the individual,	
including a child who is born to or placed for	
adoption with the individual; and	
``(C) all other individuals related by blood	
to the individual or the spouse or child described	
in subparagraph (A) or (B).	
"(16) GENETIC INFORMATION.—The term 'ge-	
netic information' means information about genes,	
gene products, or inherited characteristics that may	
derive from an individual or a family member (in-	
cluding information about a request for or receipt of	
genetic services).	

1	"(17) GENETIC SERVICES.—The term 'genetic
2	services' means health services provided to obtain, as-
3	sess, or interpret genetic information for diagnostic
4	and therapeutic purposes, and for genetic education
5	and counseling.
6	"(18) Predictive genetic information.—
7	"(A) IN GENERAL.—The term 'predictive ge-
8	netic information' means, in the absence of
9	symptoms, clinical signs, or a diagnosis of the
10	condition related to such information—
11	"(i) information about an individual's
12	genetic tests;
13	"(ii) information about genetic tests of
14	family members of the individual; or
15	"(iii) information about the occurrence
16	of a disease or disorder in family members.
17	"(B) EXCEPTIONS.—The term 'predictive
18	genetic information' shall not include—
19	"(i) information about the sex or age of
20	the individual;
21	"(ii) information derived from phys-
22	ical tests, such as the chemical, blood, or
23	urine analyses of the individual including
24	cholesterol tests; and

1	ʻʻ(iii)	information	about	physical
2	exams of the	individual.		

3 "(19) GENETIC TEST.—The term 'genetic test' 4 means the analysis of human DNA, RNA, chro-5 mosomes, proteins, and certain metabolites, including 6 analysis of genotypes, mutations, phenotypes, or 7 karyotypes, for the purpose of predicting risk of dis-8 ease in asymptomatic or undiagnosed individuals. 9 Such term does not include physical tests, such as the 10 chemical, blood, or urine analyses of the individual 11 including cholesterol tests, and physical exams of the 12 individual, in order to detect symptoms, clinical 13 signs, or a diagnosis of disease.".

(b) AMENDMENT RELATING TO THE INDIVIDUAL MARKET.—Subpart 2 of part B of title XXVII of the Public
Health Service Act, as amended by section 201, is further
amended by adding at the end the following new section: **"SEC. 2754. PROHIBITION OF HEALTH DISCRIMINATION ON**THE BASIS OF PREDICTIVE GENETIC INFOR-

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MATION.

21 "(a) PROHIBITION ON PREDICTIVE GENETIC INFORMA22 TION AS A CONDITION OF ELIGIBILITY.—A health insurance
23 issuer offering health insurance coverage in the individual
24 market may not use predictive genetic information as a
25 condition of eligibility of an individual to enroll in indi-

vidual health insurance coverage (including information
 about a request for or receipt of genetic services).

3 "(b) PROHIBITION ON PREDICTIVE GENETIC INFORMA-4 TION IN SETTING PREMIUM RATES.—A health insurance 5 issuer offering health insurance coverage in the individual 6 market shall not adjust premium rates for individuals on 7 the basis of predictive genetic information concerning such 8 an individual (including a dependent) or a family member of the individual (including information about a request 9 for or receipt of genetic services). 10

11 "(c) Collection of Predictive Genetic Informa12 TION.—

13 "(1) Limitation on requesting or requiring 14 PREDICTIVE GENETIC INFORMATION.—Except as pro-15 vided in paragraph (2), a health insurance issuer of-16 fering health insurance coverage in the individual 17 market shall not request or require predictive genetic 18 information concerning any individual (including a 19 dependent) or a family member of the individual (in-20 cluding information about a request for or receipt of 21 genetic services).

22 "(2) INFORMATION NEEDED FOR DIAGNOSIS,
23 TREATMENT, OR PAYMENT.—

24 "(A) IN GENERAL.—Notwithstanding para25 graph (1), a health insurance issuer offering

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1 health insurance coverage in the individual mar-2 ket that provides health care items and services 3 to an individual or dependent may request (but 4 may not require) that such individual or dependent disclose, or authorize the collection or 5 6 disclosure of, predictive genetic information for 7 purposes of diagnosis, treatment, or payment re-8 lating to the provision of health care items and 9 services to such individual or dependent. 10 "(B) NOTICE OF CONFIDENTIALITY PRAC-11 TICES AND DESCRIPTION OF SAFEGUARDS.—As a 12 part of a request under subparagraph (A), the 13 health insurance issuer offering health insurance 14 coverage in the individual market shall provide 15 to the individual or dependent a description of 16 the procedures in place to safeguard the con-17 fidentiality, as described in subsection (d), of 18 such predictive genetic information. 19 "(d) Confidentiality with Respect to Pre-DICTIVE GENETIC INFORMATION.— 20 21 "(1) NOTICE OF CONFIDENTIALITY PRACTICES.— 22 "(A) PREPARATION OF WRITTEN NOTICE.— 23 A health insurance issuer offering health insur-24 ance coverage in the individual market shall post 25 or provide, in writing and in a clear and con-

1	spicuous manner, notice of the issuer's confiden-
2	tiality practices, that shall include—
3	"(i) a description of an individual's
4	rights with respect to predictive genetic in-
5	formation;
6	"(ii) the procedures established by the
7	issuer for the exercise of the individual's
8	rights; and
9	"(iii) the right to obtain a copy of the
10	notice of the confidentiality practices re-
11	quired under this subsection.
12	"(B) MODEL NOTICE.—The Secretary, in
13	consultation with the National Committee on
14	Vital and Health Statistics and the National As-
15	sociation of Insurance Commissioners, and after
16	notice and opportunity for public comment, shall
17	develop and disseminate model notices of con-
18	fidentiality practices. Use of the model notice
19	shall serve as a defense against claims of receiv-
20	ing inappropriate notice.
21	"(2) ESTABLISHMENT OF SAFEGUARDS.—A
22	health insurance issuer offering health insurance cov-
23	erage in the individual market shall establish and
24	maintain appropriate administrative, technical, and
25	physical safeguards to protect the confidentiality, se-

1	curity, accuracy, and integrity of predictive genetic
2	information created, received, obtained, maintained,
3	used, transmitted, or disposed of by such issuer.".
4	(c) EFFECTIVE DATE.—The amendments made by this
5	section shall apply with respect to—
6	(1) group health plans, and health insurance
7	coverage offered in connection with group health
8	plans, for plan years beginning after 1 year after the
9	date of enactment of this Act; and
10	(2) health insurance coverage offered, sold,
11	issued, renewed, in effect, or operated in the indi-
12	vidual market after 1 year after the date of enactment
13	of this Act.
15	
14	SEC. 304. AMENDMENTS TO THE INTERNAL REVENUE CODE
14	SEC. 304. AMENDMENTS TO THE INTERNAL REVENUE CODE
14 15	SEC. 304. AMENDMENTS TO THE INTERNAL REVENUE CODE OF 1986.
14 15 16	SEC. 304. AMENDMENTS TO THE INTERNAL REVENUE CODE OF 1986. (a) PROHIBITION OF HEALTH DISCRIMINATION ON
14 15 16 17	SEC. 304. AMENDMENTS TO THE INTERNAL REVENUE CODE OF 1986. (a) Prohibition of Health Discrimination on THE BASIS OF GENETIC INFORMATION OR GENETIC SERV-
14 15 16 17 18	SEC. 304. AMENDMENTS TO THE INTERNAL REVENUE CODE OF 1986. (a) Prohibition of Health Discrimination on The Basis of Genetic Information or Genetic Serv- ICES.—
14 15 16 17 18 19	SEC. 304. AMENDMENTS TO THE INTERNAL REVENUE CODE OF 1986. (a) PROHIBITION OF HEALTH DISCRIMINATION ON THE BASIS OF GENETIC INFORMATION OR GENETIC SERV- ICES.— (1) NO ENROLLMENT RESTRICTION FOR GENETIC
 14 15 16 17 18 19 20 	SEC. 304. AMENDMENTS TO THE INTERNAL REVENUE CODE OF 1986. (a) Prohibition of Health Discrimination on The Basis of Genetic Information or Genetic Serv- ices.— (1) No enrollment restriction for genetic Services.—Section 9802(a)(1)(F) of the Internal
 14 15 16 17 18 19 20 21 	SEC. 304. AMENDMENTS TO THE INTERNAL REVENUE CODE OF 1986. (a) PROHIBITION OF HEALTH DISCRIMINATION ON THE BASIS OF GENETIC INFORMATION OR GENETIC SERV- ICES.— (1) NO ENROLLMENT RESTRICTION FOR GENETIC SERVICES.—Section 9802(a)(1)(F) of the Internal Revenue Code of 1986 is amended by inserting before
 14 15 16 17 18 19 20 21 22 	SEC. 304. AMENDMENTS TO THE INTERNAL REVENUE CODE OF 1986. (a) PROHIBITION OF HEALTH DISCRIMINATION ON THE BASIS OF GENETIC INFORMATION OR GENETIC SERV- ICES.— (1) NO ENROLLMENT RESTRICTION FOR GENETIC SERVICES.—Section 9802(a)(1)(F) of the Internal Revenue Code of 1986 is amended by inserting before the period the following: "(including information

1 (A) IN GENERAL.—Subchapter B of chapter 2 100 of the Internal Revenue Code of 1986, as amended by sections 111(b) and 201, is further 3 4 amended by adding at the end the following: 5 "SEC. 9815. PROHIBITING PREMIUM DISCRIMINATION 6 AGAINST GROUPS ON THE BASIS OF PRE-7 **DICTIVE GENETIC INFORMATION.** 8 "A group health plan shall not adjust premium or con-9 tribution amounts for a group on the basis of predictive genetic information concerning any individual (including 10 a dependent) or a family member of the individual (includ-11 ing information about a request for or receipt of genetic 12

14 (B)Conforming AMENDMENT.—Section 15 9802(b) of the Internal Revenue Code of 1986 is 16 amended by adding at the end the following: "(3) Reference to related provision.—For 17 18 a provision prohibiting the adjustment of premium or 19 contribution amounts for a group under a group 20 health plan on the basis of predictive genetic informa-21 tion (including information about a request for or the 22 receipt of genetic services), see section 9815.". 23 (C) Amendment to table of sections.— 24 The table of sections for subchapter B of chapter 25 100 of the Internal Revenue Code of 1986, as

13

services).".

1	amended by sections 111(b) and 201, is further
2	amended by adding at the end the following:
	"Sec. 9816. Prohibiting premium discrimination against groups on the basis of predictive genetic information.".
3	(b) Limitation on Collection of Predictive Ge-
4	NETIC INFORMATION.—Section 9802 of the Internal Rev-
5	enue Code of 1986 is amended by adding at the end the
6	following:
7	"(d) Collection of Predictive Genetic Informa-
8	TION.—
9	"(1) Limitation on requesting or requiring
10	PREDICTIVE GENETIC INFORMATION.—Except as pro-
11	vided in paragraph (2), a group health plan shall not
12	request or require predictive genetic information con-
13	cerning any individual (including a dependent) or a
14	family member of the individual (including informa-
15	tion about a request for or receipt of genetic services).
16	"(2) INFORMATION NEEDED FOR DIAGNOSIS,
17	TREATMENT, OR PAYMENT.—
18	"(A) IN GENERAL.—Notwithstanding para-
19	graph (1), a group health plan that provides
20	health care items and services to an individual
21	or dependent may request (but may not require)
22	that such individual or dependent disclose, or
23	authorize the collection or disclosure of, pre-
24	dictive genetic information for purposes of diag-

1	nosis, treatment, or payment relating to the pro-
2	vision of health care items and services to such
3	individual or dependent.
4	"(B) Notice of confidentiality prac-
5	TICES; DESCRIPTION OF SAFEGUARDS.—As a
6	part of a request under subparagraph (A) , the
7	group health plan shall provide to the individual
8	or dependent a description of the procedures in
9	place to safeguard the confidentiality, as de-
10	scribed in subsection (e), of such predictive ge-
11	netic information.
12	"(e) Confidentiality with Respect to Predictive
13	Genetic Information.—
14	"(1) Notice of confidentiality practices.—
15	"(A) Preparation of written notice.—
16	A group health plan shall post or provide, in
17	writing and in a clear and conspicuous manner,
18	notice of the plan's confidentiality practices, that
19	shall include—
20	"(i) a description of an individual's
21	rights with respect to predictive genetic in-
22	formation;
23	"(ii) the procedures established by the
24	plan for the exercise of the individual's
25	rights; and

"(iii) the right to obtain a copy of the
 notice of the confidentiality practices re quired under this subsection.

4 "(B) MODEL NOTICE.—The Secretary, in consultation with the National Committee on 5 6 Vital and Health Statistics and the National As-7 sociation of Insurance Commissioners, and after 8 notice and opportunity for public comment, shall 9 develop and disseminate model notices of confidentiality practices. Use of the model notice 10 11 shall serve as a defense against claims of receiv-12 ing inappropriate notice.

13 (2)Establishment OFSAFEGUARDS.—A 14 group health plan shall establish and maintain ap-15 propriate administrative, technical, and physical 16 safequards to protect the confidentiality, security, ac-17 curacy, and integrity of predictive genetic informa-18 tion created, received, obtained, maintained, used, 19 transmitted, or disposed of by such plan.".

20 (c) DEFINITIONS.—Section 9832(d) of the Internal
21 Revenue Code of 1986 is amended by adding at the end
22 the following:

23 "(6) FAMILY MEMBER.—The term 'family mem24 ber' means, with respect to an individual—
25 "(A) the spouse of the individual;

1	"(B) a dependent child of the individual,
2	including a child who is born to or placed for
3	adoption with the individual; and
4	``(C) all other individuals related by blood
5	to the individual or the spouse or child described
6	in subparagraph (A) or (B).
7	"(7) GENETIC INFORMATION.—The term 'genetic
8	information' means information about genes, gene
9	products, or inherited characteristics that may derive
10	from an individual or a family member (including
11	information about a request for or receipt of genetic
12	services).
13	"(8) GENETIC SERVICES.—The term 'genetic
14	services' means health services provided to obtain, as-
15	sess, or interpret genetic information for diagnostic
16	and therapeutic purposes, and for genetic education
17	and counseling.
18	"(9) Predictive genetic information.—
19	"(A) IN GENERAL.—The term 'predictive ge-
20	netic information' means, in the absence of
21	symptoms, clinical signs, or a diagnosis of the
22	condition related to such information—
23	"(i) information about an individual's
24	genetic tests;

1	"(ii) information about genetic tests of
2	family members of the individual; or
3	"(iii) information about the occurrence
4	of a disease or disorder in family members.
5	"(B) EXCEPTIONS.—The term 'predictive
6	genetic information' shall not include—
7	"(i) information about the sex or age of
8	the individual;
9	"(ii) information derived from phys-
10	ical tests, such as the chemical, blood, or
11	urine analyses of the individual including
12	cholesterol tests; and
13	"(iii) information about physical
14	exams of the individual.
15	"(10) GENETIC TEST.—The term 'genetic test'
16	means the analysis of human DNA, RNA, chro-
17	mosomes, proteins, and certain metabolites, including
18	analysis of genotypes, mutations, phenotypes, or
19	karyotypes, for the purpose of predicting risk of dis-
20	ease in asymptomatic or undiagnosed individuals.
21	Such term does not include physical tests, such as the
22	chemical, blood, or urine analyses of the individual
23	including cholesterol tests, and physical exams of the
24	individual, in order to detect symptoms, clinical
25	signs, or a diagnosis of disease.".

(d) EFFECTIVE DATE.—Except as provided in this sec tion, this section and the amendments made by this section
 shall apply with respect to group health plans for plan
 years beginning after 1 year after the date of the enactment
 of this Act.

6 TITLE IV—HEALTHCARE 7 RESEARCH AND QUALITY

8 SEC. 401. SHORT TITLE.

9 This title may be cited as the "Healthcare Research10 and Quality Act of 1999".

SEC. 402. AMENDMENT TO THE PUBLIC HEALTH SERVICE
 ACT.
 Title IX of the Public Health Service Act (42 U.S.C.
 299 et seq.) is amended to read as follows:
 "TITLE IX—AGENCY FOR HEALTHCARE RESEARCH AND

17 **QUALITY**

18 "PART A-ESTABLISHMENT AND GENERAL

19 **DUTIES**

20 "SEC. 901. MISSION AND DUTIES.

21 "(a) IN GENERAL.—There is established within the
22 Public Health Service an agency to be known as the Agency
23 for Healthcare Research and Quality. In carrying out this
24 subsection, the Secretary shall redesignate the Agency for

Health Care Policy and Research as the Agency for
 Healthcare Research and Quality.

3 "(b) MISSION.—The purpose of the Agency is to en-4 hance the quality, appropriateness, and effectiveness of 5 healthcare services, and access to such services, through the establishment of a broad base of scientific research and 6 7 through the promotion of improvements in clinical and 8 health system practices, including the prevention of diseases and other health conditions. The Agency shall promote 9 healthcare quality improvement by— 10

"(1) conducting and supporting research that develops and presents scientific evidence regarding all
aspects of healthcare, including—

14 "(A) the development and assessment of
15 methods for enhancing patient participation in
16 their own care and for facilitating shared pa17 tient-physician decision-making;

18 "(B) the outcomes, effectiveness, and cost-ef19 fectiveness of healthcare practices, including pre20 ventive measures and long-term care;

21 "(C) existing and innovative technologies;
22 "(D) the costs and utilization of, and access
23 to healthcare;

24 "(E) the ways in which healthcare services
25 are organized, delivered, and financed and the

1	interaction and impact of these factors on the
2	quality of patient care;
3	``(F) methods for measuring quality and
4	strategies for improving quality; and
5	"(G) ways in which patients, consumers,
6	purchasers, and practitioners acquire new infor-
7	mation about best practices and health benefits,
8	the determinants and impact of their use of this
9	information;
10	"(2) synthesizing and disseminating available
11	scientific evidence for use by patients, consumers,
12	practitioners, providers, purchasers, policy makers,
13	and educators; and
14	"(3) advancing private and public efforts to im-
15	prove healthcare quality.
16	"(c) Requirements With Respect to Rural
17	Areas and Priority Populations.—In carrying out sub-
18	section (b), the Director shall undertake and support re-
19	search, demonstration projects, and evaluations with respect
20	to the delivery of health services—
21	"(1) in rural areas (including frontier areas);
22	"(2) for low-income groups, and minority
23	groups;
24	"(3) for children;
25	"(4) for elderly; and

"(5) for people with special healthcare needs, in cluding disabilities, chronic care and end-of-life
 healthcare.

4 "(d) APPOINTMENT OF DIRECTOR.—There shall be at
5 the head of the Agency an official to be known as the Direc6 tor for Healthcare Research and Quality. The Director shall
7 be appointed by the Secretary. The Secretary, acting
8 through the Director, shall carry out the authorities and
9 duties established in this title.

10 "SEC. 902. GENERAL AUTHORITIES.

11 "(a) IN GENERAL.—In carrying out section 901(b), the 12 Director shall support demonstration projects, conduct and 13 support research, evaluations, training, research networks, 14 multi-disciplinary centers, technical assistance, and the dis-15 semination of information, on healthcare, and on systems 16 for the delivery of such care, including activities with re-17 spect to—

- 18 "(1) the quality, effectiveness, efficiency, appro19 priateness and value of healthcare services;
- 20 "(2) quality measurement and improvement;
- 21 "(3) the outcomes, cost, cost-effectiveness, and use
 22 of healthcare services and access to such services;
- 23 "(4) clinical practice, including primary care
 24 and practice-oriented research;

1	"(5) healthcare technologies, facilities, and equip-
2	ment;
3	"(6) healthcare costs, productivity, organization,
4	and market forces;
5	"(7) health promotion and disease prevention,
6	including clinical preventive services;
7	"(8) health statistics, surveys, database develop-
8	ment, and epidemiology; and
9	"(9) medical liability.
10	"(b) Health Services Training Grants.—
11	"(1) IN GENERAL.—The Director may provide
12	training grants in the field of health services research
13	related to activities authorized under subsection (a),
14	to include pre- and post-doctoral fellowships and
15	training programs, young investigator awards, and
16	other programs and activities as appropriate. In car-
17	rying out this subsection, the Director shall make use
18	of funds made available under section 487 as well as
19	other appropriated funds.
20	"(2) Requirements.—In developing priorities
21	for the allocation of training funds under this sub-
22	section, the Director shall take into consideration
23	shortages in the number of trained researchers ad-
24	dressing the priority populations.

"(c) MULTIDISCIPLINARY CENTERS.—The Director
 may provide financial assistance to assist in meeting the
 costs of planning and establishing new centers, and oper ating existing and new centers, for multidisciplinary health
 services research, demonstration projects, evaluations,
 training, and policy analysis with respect to the matters
 referred to in subsection (a).

8 "(d) Relation to Certain Authorities Regard-9 ING SOCIAL SECURITY.—Activities authorized in this sec-10 tion shall be appropriately coordinated with experiments, demonstration projects, and other related activities author-11 ized by the Social Security Act and the Social Security 12 Amendments of 1967. Activities under subsection (a)(2) of 13 this section that affect the programs under titles XVIII, XIX 14 15 and XXI of the Social Security Act shall be carried out consistent with section 1142 of such Act. 16

17 "(e) DISCLAIMER.—The Agency shall not mandate na18 tional standards of clinical practice or quality healthcare
19 standards. Recommendations resulting from projects funded
20 and published by the Agency shall include a corresponding
21 disclaimer.

22 "(f) RULE OF CONSTRUCTION.—Nothing in this sec-23 tion shall be construed to imply that the Agency's role is 24 to mandate a national standard or specific approach to 25 quality measurement and reporting. In research and quality improvement activities, the Agency shall consider a wide
 range of choices, providers, healthcare delivery systems, and
 individual preferences.

4 "PART B—HEALTHCARE IMPROVEMENT
5 RESEARCH
6 "SEC. 911. HEALTHCARE OUTCOME IMPROVEMENT RE7 SEARCH.

8 "(a) EVIDENCE RATING SYSTEMS.—In collaboration 9 with experts from the public and private sector, the Agency 10 shall identify and disseminate methods or systems that it uses to assess healthcare research results, particularly meth-11 12 ods or systems that it uses to rate the strength of the sci-13 entific evidence behind healthcare practice, recommendations in the research literature, and technology assessments. 14 15 The Agency shall make methods and systems for evidence rating widely available. Agency publications containing 16 healthcare recommendations shall indicate the level of sub-17 stantiating evidence using such methods or systems. 18

19 "(b) HEALTHCARE IMPROVEMENT RESEARCH CEN-20 TERS AND PROVIDER-BASED RESEARCH NETWORKS.—In 21 order to address the full continuum of care and outcomes 22 research, to link research to practice improvement, and to 23 speed the dissemination of research findings to community 24 practice settings, the Agency shall employ research strate-25 gies and mechanisms that will link research directly with

1	clinical practice in geographically diverse locations
2	throughout the United States, including—
3	"(1) Healthcare Improvement Research Centers
4	that combine demonstrated multidisciplinary exper-
5	tise in outcomes or quality improvement research
6	with linkages to relevant sites of care;
7	"(2) Provider-based Research Networks, includ-
8	ing plan, facility, or delivery system sites of care (es-
9	pecially primary care), that can evaluate and pro-
10	mote quality improvement; and
11	"(3) other innovative mechanisms or strategies to
12	link research with clinical practice.
13	"SEC. 912. PRIVATE-PUBLIC PARTNERSHIPS TO IMPROVE
13 14	"SEC. 912. PRIVATE-PUBLIC PARTNERSHIPS TO IMPROVE ORGANIZATION AND DELIVERY.
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14	ORGANIZATION AND DELIVERY.
14 15	ORGANIZATION AND DELIVERY. "(a) Support for Efforts To Develop Informa-
14 15 16	ORGANIZATION AND DELIVERY. "(a) Support for Efforts To Develop Informa- tion on Quality.—
14 15 16 17	ORGANIZATION AND DELIVERY. "(a) Support for Efforts To Develop Informa- tion on Quality.— "(1) Scientific and technical support.—In
14 15 16 17 18	ORGANIZATION AND DELIVERY. "(a) Support for Efforts To Develop Informa- TION ON QUALITY.— "(1) Scientific and technical support.—In its role as the principal agency for healthcare research
 14 15 16 17 18 19 	ORGANIZATION AND DELIVERY. "(a) SUPPORT FOR EFFORTS TO DEVELOP INFORMA- TION ON QUALITY.— "(1) SCIENTIFIC AND TECHNICAL SUPPORT.—In its role as the principal agency for healthcare research and quality, the Agency may provide scientific and
 14 15 16 17 18 19 20 	ORGANIZATION AND DELIVERY. "(a) SUPPORT FOR EFFORTS TO DEVELOP INFORMA- TION ON QUALITY.— "(1) SCIENTIFIC AND TECHNICAL SUPPORT.—In its role as the principal agency for healthcare research and quality, the Agency may provide scientific and technical support for private and public efforts to im-
 14 15 16 17 18 19 20 21 	ORGANIZATION AND DELIVERY. "(a) SUPPORT FOR EFFORTS TO DEVELOP INFORMA- TION ON QUALITY.— "(1) SCIENTIFIC AND TECHNICAL SUPPORT.—In its role as the principal agency for healthcare research and quality, the Agency may provide scientific and technical support for private and public efforts to im- prove healthcare quality, including the activities of

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1	``(A) the identification and assessment of
2	methods for the evaluation of the health of—
3	"(i) enrollees in health plans by type of
4	plan, provider, and provider arrangements;
5	and
6	"(ii) other populations, including those
7	receiving long-term care services;
8	``(B) the ongoing development, testing, and
9	dissemination of quality measures, including
10	measures of health and functional outcomes;
11	(C) the compilation and dissemination of
12	healthcare quality measures developed in the pri-
13	vate and public sector;
14	``(D) assistance in the development of im-
15	proved healthcare information systems;
16	((E) the development of survey tools for the
17	purpose of measuring participant and bene-
18	ficiary assessments of their healthcare; and
19	``(F) identifying and disseminating infor-
20	mation on mechanisms for the integration of in-
21	formation on quality into purchaser and con-
22	sumer decision-making processes.
23	"(b) Centers for Education and Research on
24	Therapeutics —

24 THERAPEUTICS.—

1	"(1) IN GENERAL.—The Secretary, acting
2	through the Director and in consultation with the
3	Commissioner of Food and Drugs, shall establish a
4	program for the purpose of making one or more
5	grants for the establishment and operation of one or
6	more centers to carry out the activities specified in
7	paragraph (2).
8	"(2) REQUIRED ACTIVITIES.—The activities re-
9	ferred to in this paragraph are the following:
10	"(A) The conduct of state-of-the-art clinical,
11	laboratory, or health services research for the fol-
12	lowing purposes:
13	"(i) To increase awareness of—
14	((I) new uses of drugs, biological
15	products, and devices;
16	"(II) ways to improve the effective
17	use of drugs, biological products, and
18	devices; and
19	"(III) risks of new uses and risks
20	of combinations of drugs and biological
21	products.
22	"(ii) To provide objective clinical in-
23	formation to the following individuals and
24	entities:

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1	"(I) Healthcare practitioners and
2	other providers of healthcare goods or
3	services.
4	"(II) Pharmacists, pharmacy ben-
5	efit managers and purchasers.
6	"(III) Health maintenance orga-
7	nizations and other managed
8	healthcare organizations.
9	"(IV) Healthcare insurers and
10	governmental agencies.
11	"(V) Patients and consumers.
12	"(iii) To improve the quality of
13	healthcare while reducing the cost of
14	Healthcare through—
15	"(I) an increase in the appro-
16	priate use of drugs, biological products,
17	or devices; and
18	"(II) the prevention of adverse ef-
19	fects of drugs, biological products, and
20	devices and the consequences of such ef-
21	fects, such as unnecessary hospitaliza-
22	tions.
23	(B) The conduct of research on the com-
24	parative effectiveness, cost-effectiveness, and safe-
25	ty of drugs, biological products, and devices.

1	"(C) Such other activities as the Secretary
2	determines to be appropriate, except that grant
3	funds may not be used by the Secretary in con-
4	ducting regulatory review of new drugs.
5	"(c) Reducing Errors in Medicine.—The Director
6	shall conduct and support research and build private-public
7	partnerships to—
8	"(1) identify the causes of preventable healthcare
9	errors and patient injury in healthcare delivery;
10	"(2) develop, demonstrate, and evaluate strate-
11	gies for reducing errors and improving patient safety;
12	and
13	"(3) promote the implementation of effective
14	strategies throughout the healthcare industry.
15	"SEC. 913. INFORMATION ON QUALITY AND COST OF CARE.
16	"(a) IN GENERAL.—In carrying out 902(a), the Direc-
17	tor shall—
18	"(1) conduct a survey to collect data on a na-
19	tionally representative sample of the population on
20	the cost, use and, for fiscal year 2001 and subsequent
21	fiscal years, quality of healthcare, including the types
22	of healthcare services Americans use, their access to
23	healthcare services, frequency of use, how much is
24	paid for the services used, the source of those pay-
25	ments, the types and costs of private health insurance,

1	access, satisfaction, and quality of care for the general
2	population including rural residents and for the pop-
3	ulations identified in section 901(c); and
4	"(2) develop databases and tools that provide in-
5	formation to States on the quality, access, and use of
6	healthcare services provided to their residents.
7	"(b) Quality and Outcomes Information.—
8	"(1) In GENERAL.—Beginning in fiscal year
9	2001, the Director shall ensure that the survey con-
10	ducted under subsection (a)(1) will—
11	(A) identify determinants of health out-
12	comes and functional status, and their relation-
13	ships to healthcare access and use, determine the
14	ways and extent to which the priority popu-
15	lations enumerated in section 901(c) differ from
16	the general population with respect to such vari-
17	ables, measure changes over time with respect to
18	such variable, and monitor the overall national
19	impact of changes in Federal and State policy
20	on healthcare;
21	(B) provide information on the quality of
22	care and patient outcomes for frequently occur-
23	ring clinical conditions for a nationally rep-
24	resentative sample of the population including
25	rural residents; and

1	(C) provide reliable national estimates for
2	children and persons with special healthcare
3	needs through the use of supplements or periodic
4	expansions of the survey.
5	In expanding the Medical Expenditure Panel Survey,
6	as in existence on the date of enactment of this title,
7	in fiscal year 2001 to collect information on the qual-
8	ity of care, the Director shall take into account any
9	outcomes measurements generally collected by private
10	sector accreditation organizations.
11	"(2) ANNUAL REPORT.—Beginning in fiscal year
12	2003, the Secretary, acting through the Director, shall
13	submit to Congress an annual report on national
14	trends in the quality of healthcare provided to the
15	American people.
16	"SEC. 914. INFORMATION SYSTEMS FOR HEALTHCARE IM-
17	PROVEMENT.
18	"(a) IN GENERAL.—In order to foster a range of inno-
19	vative approaches to the management and communication
20	of health information, the Agency shall support research,
21	evaluations and initiatives to advance—

"(1) the use of information systems for the study
of healthcare quality, including the generation of both
individual provider and plan-level comparative performance data;

1	"(2) training for healthcare practitioners and re-
2	searchers in the use of information systems;
3	"(3) the creation of effective linkages between
4	various sources of health information, including the
5	development of information networks;
6	"(4) the delivery and coordination of evidence-
7	based healthcare services, including the use of real-
8	time healthcare decision-support programs;
9	"(5) the utility and comparability of health in-
10	formation data and medical vocabularies by address-
11	ing issues related to the content, structure, definitions
12	and coding of such information and data in consulta-
13	tion with appropriate Federal, State and private en-
14	tities;
15	"(6) the use of computer-based health records in
16	all settings for the development of personal health
17	records for individual health assessment and mainte-
18	nance, and for monitoring public health and outcomes
19	of care within populations; and
20	"(7) the protection of individually identifiable
21	information in health services research and healthcare
22	quality improvement.
23	"(b) Demonstration.—The Agency shall support
24	demonstrations into the use of new information tools aimed

3 "SEC. 915. RESEARCH SUPPORTING PRIMARY CARE AND AC-4 **CESS IN UNDERSERVED AREAS.** "(a) Preventive Services Task Force.— 5 6 "(1) Establishment and purpose.—The Di-7 rector may periodically convene a Preventive Services 8 Task Force to be composed of individuals with appro-9 priate expertise. Such a task force shall review the 10 scientific evidence related to the effectiveness, appro-11 priateness, and cost-effectiveness of clinical preventive 12 services for the purpose of developing recommenda-13 tions for the healthcare community, and updating 14 previous clinical preventive recommendations. 15 "(2) ROLE OF AGENCY.—The Agency shall pro-16 vide ongoing administrative, research, and technical 17 support for the operations of the Preventive Services 18 Task Force, including coordinating and supporting 19 the dissemination of the recommendations of the Task 20 Force. 21 "(3) OPERATION.—In carrying out its respon-22 sibilities under paragraph (1), the Task Force is not 23 subject to the provisions of Appendix 2 of title 5,

24 United States Code.

25 "(b) PRIMARY CARE RESEARCH.—

1 "(1) IN GENERAL.—There is established within 2 the Agency a Center for Primary Care Research (re-3 ferred to in this subsection as the 'Center') that shall 4 serve as the principal source of funding for primary 5 care practice research in the Department of Health 6 and Human Services. For purposes of this paragraph, 7 primary care research focuses on the first contact 8 when illness or health concerns arise, the diagnosis, 9 treatment or referral to specialty care, preventive 10 care, and the relationship between the clinician and 11 the patient in the context of the family and commu-12 nity. 13 "(2) RESEARCH.—In carrying out this section, 14 Center shall conduct and support research the 15 concerning— "(A) the nature and characteristics of pri-16 17 mary care practice; 18 (B) the management of commonly occur-19 ring clinical problems; (C) the management of undifferentiated 20 21 clinical problems; and "(D) the continuity and coordination of 22 23 health services.

1 "SEC. 916. CLINICAL PRACTICE AND TECHNOLOGY INNOVA-

2	TION.
3	"(a) IN GENERAL.—The Director shall promote inno-
4	vation in evidence-based clinical practice and healthcare
5	technologies by—
6	((1) conducting and supporting research on the
7	development, diffusion, and use of healthcare tech-
8	nology;
9	"(2) developing, evaluating, and disseminating
10	methodologies for assessments of healthcare practices
11	and healthcare technologies;
12	((3) conducting intramural and supporting ex-
13	tramural assessments of existing and new healthcare
14	practices and technologies;
15	"(4) promoting education, training, and pro-
16	viding technical assistance in the use of healthcare
17	practice and healthcare technology assessment meth-
18	odologies and results; and
19	"(5) working with the National Library of Medi-
20	cine and the public and private sector to develop an
21	electronic clearinghouse of currently available assess-
22	ments and those in progress.
23	"(b) Specification of Process.—
24	"(1) IN GENERAL.—Not later than December 31,
25	2000, the Director shall develop and publish a de-
26	scription of the methodology used by the Agency and
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its contractors in conducting practice and technology assessment.

3	"(2) CONSULTATIONS.—In carrying out this sub-
4	section, the Director shall cooperate and consult with
5	the Assistant Secretary for Health, the Administrator
б	of the Health Care Financing Administration, the Di-
7	rector of the National Institutes of Health, the Com-
8	missioner of Food and Drugs, and the heads of any
9	other interested Federal department or agency, and
10	shall seek input, where appropriate, from professional
11	societies and other private and public entities.
12	"(3) Methodology.—The Director, in devel-
13	oping assessment methodology, shall consider—
14	"(A) safety, efficacy, and effectiveness;
15	"(B) legal, social, and ethical implications;
16	"(C) costs, benefits, and cost-effectiveness;
17	(D) comparisons to alternate technologies
18	and practices; and
19	((E) requirements of Food and Drug Ad-
20	ministration approval to avoid duplication.
21	"(c) Specific Assessments.—
22	"(1) IN GENERAL.—The Director shall conduct
23	or support specific assessments of healthcare tech-
24	nologies and practices.

1	"(2) Requests for assessments.—The Direc-
2	tor is authorized to conduct or support assessments,
3	on a reimbursable basis, for the Health Care Financ-
4	ing Administration, the Department of Defense, the
5	Department of Veterans Affairs, the Office of Per-
6	sonnel Management, and other public or private enti-
7	ties.
8	"(3) GRANTS AND CONTRACTS.—In addition to
9	conducting assessments, the Director may make
10	grants to, or enter into cooperative agreements or con-
11	tracts with, entities described in paragraph (4) for
12	the purpose of conducting assessments of experi-

mental, emerging, existing, or potentially outmoded
healthcare technologies, and for related activities.

"(4) ELIGIBLE ENTITIES.—An entity described 15 in this paragraph is an entity that is determined to 16 17 be appropriate by the Director, including academic 18 medical centers, research institutions and organiza-19 tions, professional organizations, third party payers, 20 governmental agencies, and consortia of appropriate 21 research entities established for the purpose of con-22 ducting technology assessments.

23 "SEC. 917. COORDINATION OF FEDERAL GOVERNMENT

- 24
- QUALITY IMPROVEMENT EFFORTS.
- 25 "(a) REQUIREMENT.—

1	"(1) IN GENERAL.—To avoid duplication and
2	ensure that Federal resources are used efficiently and
3	effectively, the Secretary, acting through the Director,
4	shall coordinate all research, evaluations, and dem-
5	onstrations related to health services research, quality
6	measurement and quality improvement activities un-
7	dertaken and supported by the Federal Government.
8	"(2) Specific activities.—The Director, in col-
9	laboration with the appropriate Federal officials rep-
10	resenting all concerned executive agencies and depart-
11	ments, shall develop and manage a process to—
12	"(A) improve interagency coordination, pri-
13	ority setting, and the use and sharing of research
14	findings and data pertaining to Federal quality
15	improvement programs, technology assessment,
16	and health services research;
17	``(B) strengthen the research information
18	infrastructure, including databases, pertaining
19	to Federal health services research and healthcare
20	quality improvement initiatives;
21	``(C) set specific goals for participating
22	agencies and departments to further health serv-
23	ices research and healthcare quality improve-
24	ment; and

1	``(D) strengthen the management of Federal
2	healthcare quality improvement programs.
3	"(b) Study by the Institute of Medicine.—
4	"(1) IN GENERAL.—To provide Congress, the De-
5	partment of Health and Human Services, and other
6	relevant departments with an independent, external
7	review of their quality oversight, quality improvement
8	and quality research programs, the Secretary shall
9	enter into a contract with the Institute of Medicine—
10	"(A) to describe and evaluate current qual-
11	ity improvement, quality research and quality
12	monitoring processes through—
13	"(i) an overview of pertinent health
14	services research activities and quality im-
15	provement efforts conducted by all Federal
16	programs, with particular attention paid to
17	those under titles XVIII, XIX, and XXI of
18	the Social Security Act; and
19	"(ii) a summary of the partnerships
20	that the Department of Health and Human
21	Services has pursued with private accredi-
22	tation, quality measurement and improve-
23	ment organizations; and
24	``(B) to identify options and make rec-
25	ommendations to improve the efficiency and ef-

1	fectiveness of quality improvement programs
2	through—
3	"(i) the improved coordination of ac-
4	tivities across the medicare, medicaid and
5	child health insurance programs under titles
6	XVIII, XIX and XXI of the Social Security
7	Act and health services research programs;
8	"(ii) the strengthening of patient choice
9	and participation by incorporating state-of-
10	the-art quality monitoring tools and mak-
11	ing information on quality available; and
12	"(iii) the enhancement of the most ef-
13	fective programs, consolidation as appro-
14	priate, and elimination of duplicative ac-
15	tivities within various federal agencies.
16	"(2) Requirements.—
17	"(A) IN GENERAL.—The Secretary shall
18	enter into a contract with the Institute of Medi-
19	cine for the preparation—
20	"(i) not later than 12 months after the
21	date of enactment of this title, of a report
22	providing an overview of the quality im-
23	provement programs of the Department of
24	Health and Human Services for the medi-
25	care, medicaid, and CHIP programs under
1	titles XVIII, XIX, and XXI of the Social Se-
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2	curity Act; and
3	"(ii) not later than 24 months after the
4	date of enactment of this title, of a final re-
5	port containing recommendations.
6	"(B) REPORTS.—The Secretary shall sub-
7	mit the reports described in subparagraph (A) to
8	the Committee on Finance and the Committee on
9	Health, Education, Labor, and Pensions of the
10	Senate and the Committee on Ways and Means
11	and the Committee on Commerce of the House of
12	Representatives.
13	"PART C—GENERAL PROVISIONS
14	"SEC. 921. ADVISORY COUNCIL FOR HEALTHCARE RE-
15	SEARCH AND QUALITY.
16	"(a) ESTABLISHMENT.—There is established an advi-
17	sory council to be known as the Advisory Council for
18	Healthcare Research and Quality.
19	"(b) DUTIES.—
20	"(1) IN GENERAL.—The Advisory Council shall
21	advise the Secretary and the Director with respect to
22	activities proposed or undertaken to carry out the
23	purpose of the Agency under section 901(b).
24	"(2) CERTAIN RECOMMENDATIONS.—Activities of
25	the Advisory Council under paragraph (1) shall in-

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1	clude making recommendations to the Director
2	regarding—
3	"(A) priorities regarding healthcare re-
4	search, especially studies related to quality, out-
5	comes, cost and the utilization of, and access to,
6	healthcare services;
7	(B) the field of healthcare research and re-
8	lated disciplines, especially issues related to
9	training needs, and dissemination of informa-
10	tion pertaining to healthcare quality; and
11	"(C) the appropriate role of the Agency in
12	each of these areas in light of private sector ac-
13	tivity and identification of opportunities for
14	public-private sector partnerships.
15	"(c) Membership.—
16	"(1) IN GENERAL.—The Advisory Council shall,
17	in accordance with this subsection, be composed of ap-
18	pointed members and ex officio members. All members
19	of the Advisory Council shall be voting members other
20	than the individuals designated under paragraph
21	(3)(B) as ex officio members.
22	"(2) APPOINTED MEMBERS.—The Secretary shall
23	appoint to the Advisory Council 21 appropriately
24	qualified individuals. At least 17 members of the Ad-
25	visory Council shall be representatives of the public

1	who are not officers or employees of the United States.
2	The Secretary shall ensure that the appointed mem-
3	bers of the Council, as a group, are representative of
4	professions and entities concerned with, or affected by,
5	activities under this title and under section 1142 of
6	the Social Security Act. Of such members—
7	``(A) 4 shall be individuals distinguished in
8	the conduct of research, demonstration projects,
9	and evaluations with respect to healthcare;
10	``(B) 4 shall be individuals distinguished in
11	the practice of medicine of which at least 1 shall
12	be a primary care practitioner;
13	``(C) 3 shall be individuals distinguished in
14	the other health professions;
15	``(D) 4 shall be individuals either rep-
16	resenting the private healthcare sector, including
17	health plans, providers, and purchasers or indi-
18	viduals distinguished as administrators of
19	healthcare delivery systems;
20	``(E) 4 shall be individuals distinguished in
21	the fields of healthcare quality improvement, eco-
22	nomics, information systems, law, ethics, busi-
23	ness, or public policy, including at least 1 indi-
24	vidual specializing in rural aspects in 1 or more
25	of these fields; and

1	((F) 2 shall be individuals representing the)
2	interests of patients and consumers of healthcare.
3	"(3) Ex officio members.—The Secretary shall
4	designate as ex officio members of the Advisory
5	Council—
6	"(A) the Assistant Secretary for Health, the
7	Director of the National Institutes of Health, the
8	Director of the Centers for Disease Control and
9	Prevention, the Administrator of the Health Care
10	Financing Administration, the Assistant Sec-
11	retary of Defense (Health Affairs), and the
12	Under Secretary for Health of the Department of
13	Veterans Affairs; and
14	"(B) such other Federal officials as the Sec-
15	retary may consider appropriate.
16	"(d) TERMS.—Members of the Advisory Council ap-
17	pointed under subsection $(c)(2)$ shall serve for a term of 3
18	years. A member of the Council appointed under such sub-
19	section may continue to serve after the expiration of the
20	term of the members until a successor is appointed.
21	"(e) VACANCIES.—If a member of the Advisory Council
22	appointed under subsection $(c)(2)$ does not serve the full
23	term applicable under subsection (d), the individual ap-
24	pointed to fill the resulting vacancy shall be appointed for

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the remainder of the term of the predecessor of the indi vidual.

3 "(f) CHAIR.—The Director shall, from among the
4 members of the Advisory Council appointed under sub5 section (c)(2), designate an individual to serve as the chair
6 of the Advisory Council.

7 "(g) MEETINGS.—The Advisory Council shall meet not
8 less than once during each discrete 4-month period and
9 shall otherwise meet at the call of the Director or the chair.
10 "(h) COMPENSATION AND REIMBURSEMENT OF EX11 PENSES.—

12 "(1) APPOINTED MEMBERS.—Members of the Ad-13 visory Council appointed under subsection (c)(2) shall 14 receive compensation for each day (including travel 15 time) engaged in carrying out the duties of the Advi-16 sory Council unless declined by the member. Such 17 compensation may not be in an amount in excess of 18 the daily equivalent of the annual rate of basic pay 19 prescribed for level IV of the Executive Schedule 20 under section 5315 of title 5, United States Code, for 21 each day during which such member is engaged in the 22 performance of the duties of the Advisory Council.

23 "(2) EX OFFICIO MEMBERS.—Officials des24 ignated under subsection (c)(3) as ex officio members
25 of the Advisory Council may not receive compensation

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1	for service on the Advisory Council in addition to the
2	compensation otherwise received for duties carried out
3	as officers of the United States.
4	"(i) STAFF.—The Director shall provide to the Advi-
5	sory Council such staff, information, and other assistance
6	as may be necessary to carry out the duties of the Council.
7	"SEC. 922. PEER REVIEW WITH RESPECT TO GRANTS AND
8	CONTRACTS.
9	"(a) Requirement of Review.—
10	"(1) IN GENERAL.—Appropriate technical and
11	scientific peer review shall be conducted with respect
12	to each application for a grant, cooperative agree-
13	ment, or contract under this title.
14	"(2) Reports to director.—Each peer review
15	group to which an application is submitted pursuant
16	to paragraph (1) shall report its finding and rec-
17	ommendations respecting the application to the Direc-
18	tor in such form and in such manner as the Director
19	shall require.
20	"(b) Approval as Precondition of Awards.—The
21	Director may not approve an application described in sub-
22	section $(a)(1)$ unless the application is recommended for ap-
23	proval by a peer review group established under subsection
24	(c).
25	"(c) Establishment of Peer Review Groups —

25 "(c) Establishment of Peer Review Groups.—

1	"(1) IN GENERAL.—The Director shall establish
2	such technical and scientific peer review groups as
3	may be necessary to carry out this section. Such
4	groups shall be established without regard to the pro-
5	visions of title 5, United States Code, that govern ap-
6	pointments in the competitive service, and without re-
7	gard to the provisions of chapter 51, and subchapter
8	III of chapter 53, of such title that relate to classifica-
9	tion and pay rates under the General Schedule.
10	"(2) Membership.—The members of any peer
11	review group established under this section shall be
12	appointed from among individuals who by virtue of
13	their training or experience are eminently qualified
14	to carry out the duties of such peer review group. Of-
15	ficers and employees of the United States may not
16	constitute more than 25 percent of the membership of
17	any such group. Such officers and employees may not
18	receive compensation for service on such groups in ad-
19	dition to the compensation otherwise received for these
20	duties carried out as such officers and employees.
21	"(3) DURATION.—Notwithstanding section $14(a)$
22	of the Federal Advisory Committee Act, peer review
23	groups established under this section may continue in
24	existence until otherwise provided by law.

"(4) QUALIFICATIONS.—Members of any peer-review group shall, at a minimum, meet the following requirements:

4 "(A) Such members shall agree in writing
5 to treat information received, pursuant to their
6 work for the group, as confidential information,
7 except that this subparagraph shall not apply to
8 public records and public information.

9 "(B) Such members shall agree in writing 10 to recuse themselves from participation in the 11 peer-review of specific applications which present 12 a potential personal conflict of interest or ap-13 pearance of such conflict, including employment 14 in a directly affected organization, stock owner-15 ship, or any financial or other arrangement that 16 might introduce bias in the process of peer-re-17 view.

18 "(d) Authority for Procedural Adjustments in 19 CERTAIN CASES.—In the case of applications for financial 20 assistance whose direct costs will not exceed \$100,000, the 21 Director may make appropriate adjustments in the proce-22 dures otherwise established by the Director for the conduct 23 of peer review under this section. Such adjustments may 24 be made for the purpose of encouraging the entry of individuals into the field of research, for the purpose of encour-25

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1 aging clinical practice-oriented or provider-based research, and for such other purposes as the Director may determine 2 3 to be appropriate. 4 "(e) REGULATIONS.—The Director shall issue regula-5 tions for the conduct of peer review under this section. 6 **"SEC. 923. CERTAIN PROVISIONS WITH RESPECT TO DEVEL-**7 **OPMENT, COLLECTION, AND DISSEMINATION** 8 OF DATA. 9 "(a) Standards With Respect to Utility of 10 DATA.— 11 "(1) IN GENERAL.—To ensure the utility, accu-12 racy, and sufficiency of data collected by or for the 13 Agency for the purpose described in section 901(b), 14 the Director shall establish standard methods for de-15 veloping and collecting such data, taking into consideration— 16 17 "(A) other Federal health data collection 18 standards; and 19 (B) the differences between typesof20 healthcare plans, delivery systems, healthcare 21 providers, and provider arrangements. 22 "(2) Relationship with other department 23 PROGRAMS.—In any case where standards under 24 paragraph (1) may affect the administration of other 25 programs carried out by the Department of Health

1	and Human Services, including the programs under
2	title XVIII, XIX or XXI of the Social Security Act,
3	or may affect health information that is subject to a
4	standard developed under part C of title XI of the So-
5	cial Security Act, they shall be in the form of rec-
6	ommendations to the Secretary for such program.
7	"(b) Statistics and Analyses.—The Director
8	shall—
9	"(1) take appropriate action to ensure that sta-
10	tistics and analyses developed under this title are of
11	high quality, timely, and duly comprehensive, and
12	that the statistics are specific, standardized, and ade-
13	quately analyzed and indexed; and
14	"(2) publish, make available, and disseminate
15	such statistics and analyses on as wide a basis as is
16	practicable.
17	"(c) Authority Regarding Certain Requests.—
18	Upon request of a public or private entity, the Director may
19	conduct or support research or analyses otherwise author-
20	ized by this title pursuant to arrangements under which
21	such entity will pay the cost of the services provided.
22	Amounts received by the Director under such arrangements
23	shall be available to the Director for obligation until ex-
24	pended.

2 "(a) IN GENERAL.—The Director shall—

3 "(1) without regard to section 501 of title 44,
4 United States Code, promptly publish, make avail5 able, and otherwise disseminate, in a form under6 standable and on as broad a basis as practicable so
7 as to maximize its use, the results of research, dem8 onstration projects, and evaluations conducted or sup9 ported under this title;

"(2) ensure that information disseminated by the
Agency is science-based and objective and undertakes
consultation as necessary to assess the appropriateness and usefulness of the presentation of information
that is targeted to specific audiences;

15 "(3) promptly make available to the public data
16 developed in such research, demonstration projects,
17 and evaluations;

18 "(4) provide, in collaboration with the National 19 Library of Medicine where appropriate, indexing, ab-20 stracting, translating, publishing, and other services 21 leading to a more effective and timely dissemination 22 of information on research, demonstration projects, 23 and evaluations with respect to healthcare to public 24 and private entities and individuals engaged in the 25 improvement of healthcare delivery and the general 26 public, and undertake programs to develop new or

1	mproved methods for making such information avail	!-
2	uble; and	

3 "(5) as appropriate, provide technical assistance
4 to State and local government and health agencies
5 and conduct liaison activities to such agencies to fos6 ter dissemination.

7 "(b) PROHIBITION AGAINST RESTRICTIONS.—Except
8 as provided in subsection (c), the Director may not restrict
9 the publication or dissemination of data from, or the results
10 of, projects conducted or supported under this title.

11 "(c) Limitation on Use of Certain Informa-12 TION.—No information, if an establishment or person sup-13 plying the information or described in it is identifiable, obtained in the course of activities undertaken or supported 14 15 under this title may be used for any purpose other than the purpose for which it was supplied unless such establish-16 ment or person has consented (as determined under regula-17 tions of the Director) to its use for such other purpose. Such 18 information may not be published or released in other form 19 if the person who supplied the information or who is de-20 21 scribed in it is identifiable unless such person has consented 22 (as determined under regulations of the Director) to its pub-23 lication or release in other form.

24 "(d) PENALTY.—Any person who violates subsection
25 (c) shall be subject to a civil monetary penalty of not more

than \$10,000 for each such violation involved. Such penalty
 shall be imposed and collected in the same manner as civil
 money penalties under subsection (a) of section 1128A of
 the Social Security Act are imposed and collected.

5 "SEC. 925. ADDITIONAL PROVISIONS WITH RESPECT TO 6 GRANTS AND CONTRACTS.

7 "(a) FINANCIAL CONFLICTS OF INTEREST.—With re8 spect to projects for which awards of grants, cooperative
9 agreements, or contracts are authorized to be made under
10 this title, the Director shall by regulation define—

11 "(1) the specific circumstances that constitute fi-12 nancial interests in such projects that will, or may be 13 reasonably expected to, create a bias in favor of ob-14 taining results in the projects that are consistent with 15 such interests; and

16 "(2) the actions that will be taken by the Direc17 tor in response to any such interests identified by the
18 Director.

19 "(b) REQUIREMENT OF APPLICATION.—The Director 20 may not, with respect to any program under this title au-21 thorizing the provision of grants, cooperative agreements, 22 or contracts, provide any such financial assistance unless 23 an application for the assistance is submitted to the Sec-24 retary and the application is in such form, is made in such 25 manner, and contains such agreements, assurances, and information as the Director determines to be necessary to
 carry out the program in involved.

3 "(c) PROVISION OF SUPPLIES AND SERVICES IN LIEU
4 OF FUNDS.—

"(1) IN GENERAL.—Upon the request of an enti-5 6 ty receiving a grant, cooperative agreement, or contract under this title, the Secretary may, subject to 7 8 paragraph (2), provide supplies, equipment, and serv-9 ices for the purpose of aiding the entity in carrying 10 out the project involved and, for such purpose, may 11 detail to the entity any officer or employee of the De-12 partment of Health and Human Services.

13 "(2) Corresponding reduction in funds.— 14 With respect to a request described in paragraph (1). 15 the Secretary shall reduce the amount of the financial 16 assistance involved by an amount equal to the costs 17 of detailing personnel and the fair market value of 18 any supplies, equipment, or services provided by the 19 Director. The Secretary shall, for the payment of ex-20 penses incurred in complying with such request, ex-21 pend the amounts withheld.

"(d) APPLICABILITY OF CERTAIN PROVISIONS WITH
RESPECT TO CONTRACTS.—Contracts may be entered into
under this part without regard to sections 3648 and 3709
of the Revised Statutes (31 U.S.C. 529; 41 U.S.C. 5).

1 "SEC. 926. CERTAIN ADMINISTRATIVE AUTHORITIES.

2 "(a) Deputy Director and Other Officers and
3 Employees.—

4 "(1) DEPUTY DIRECTOR.—The Director may ap5 point a deputy director for the Agency.

6 "(2) OTHER OFFICERS AND EMPLOYEES.—The 7 Director may appoint and fix the compensation of 8 such officers and employees as may be necessary to 9 carry out this title. Except as otherwise provided by 10 law, such officers and employees shall be appointed in 11 accordance with the civil service laws and their com-12 pensation fixed in accordance with title 5, United 13 States Code.

14 "(b) FACILITIES.—The Secretary, in carrying out this
15 title—

16 "(1) may acquire, without regard to the Act of 17 March 3, 1877 (40 U.S.C. 34), by lease or otherwise 18 through the Director of General Services, buildings or 19 portions of buildings in the District of Columbia or 20 communities located adjacent to the District of Co-21 lumbia for use for a period not to exceed 10 years; 22 and

23 "(2) may acquire, construct, improve, repair, op24 erate, and maintain laboratory, research, and other
25 necessary facilities and equipment, and such other

real or personal property (including patents) as the
 Secretary deems necessary.

3 "(c) PROVISION OF FINANCIAL ASSISTANCE.—The Di4 rector, in carrying out this title, may make grants to public
5 and nonprofit entities and individuals, and may enter into
6 cooperative agreements or contracts with public and private
7 entities and individuals.

8 "(d) UTILIZATION OF CERTAIN PERSONNEL AND RE9 SOURCES.—

10 "(1) Department of health and human 11 SERVICES.—The Director, in carrying out this title, 12 may utilize personnel and equipment, facilities, and 13 other physical resources of the Department of Health 14 and Human Services, permit appropriate (as deter-15 mined by the Secretary) entities and individuals to 16 utilize the physical resources of such Department, and 17 provide technical assistance and advice.

"(2) OTHER AGENCIES.—The Director, in carrying out this title, may use, with their consent, the
services, equipment, personnel, information, and facilities of other Federal, State, or local public agencies, or of any foreign government, with or without
reimbursement of such agencies.

24 "(e) CONSULTANTS.—The Secretary, in carrying out
25 this title, may secure, from time to time and for such peri-

ods as the Director deems advisable but in accordance with
 section 3109 of title 5, United States Code, the assistance
 and advice of consultants from the United States or abroad.
 "(f) EXPERTS.—

"(1) IN GENERAL.—The Secretary may, in car-5 6 rying out this title, obtain the services of not more 7 than 50 experts or consultants who have appropriate 8 scientific or professional qualifications. Such experts or consultants shall be obtained in accordance with 9 section 3109 of title 5, United States Code, except that 10 11 the limitation in such section on the duration of serv-12 ice shall not apply.

13 "(2) TRAVEL EXPENSES.—

14 "(A) IN GENERAL.—Experts and consult15 ants whose services are obtained under para16 graph (1) shall be paid or reimbursed for their
17 expenses associated with traveling to and from
18 their assignment location in accordance with sec19 tions 5724, 5724a(a), 5724a(c), and 5726(C) of
20 title 5, United States Code.

21 "(B) LIMITATION.—Expenses specified in
22 subparagraph (A) may not be allowed in connec23 tion with the assignment of an expert or consult24 ant whose services are obtained under paragraph
25 (1) unless and until the expert agrees in writing

1 to complete the entire period of assignment, or 1 2 year, whichever is shorter, unless separated or re-3 assigned for reasons that are beyond the control 4 of the expert or consultant and that are acceptable to the Secretary. If the expert or consultant 5 6 violates the agreement, the money spent by the United States for the expenses specified in sub-7 8 paragraph (A) is recoverable from the expert or 9 consultant as a statutory obligation oved to the 10 United States. The Secretary may waive in 11 whole or in part a right of recovery under this 12 subparagraph.

13 "(g) VOLUNTARY AND UNCOMPENSATED SERVICES.—
14 The Director, in carrying out this title, may accept vol15 untary and uncompensated services.

16 "SEC. 927. FUNDING.

17 "(a) INTENT.—To ensure that the United States's investment in biomedical research is rapidly translated into 18 improvements in the quality of patient care, there must be 19 a corresponding investment in research on the most effective 20 21 clinical and organizational strategies for use of these find-22 ings in daily practice. The authorization levels in sub-23 section (b) provide for a proportionate increase in healthcare research as the United States investment in bio-24 medical research increases. 25

"(b) AUTHORIZATION OF APPROPRIATIONS.—For the
 purpose of carrying out this title, there are authorized to
 be appropriated \$250,000,000 for fiscal year 2000, and such
 sums as may be necessary for each of the fiscal years 2001
 through 2006.

6 "(c) EVALUATIONS.—In addition to amounts available 7 pursuant to subsection (b) for carrying out this title, there 8 shall be made available for such purpose, from the amounts 9 made available pursuant to section 241 (relating to evalua-10 tions), an amount equal to 40 percent of the maximum 11 amount authorized in such section 241 to be made available 12 for a fiscal year.

13 "SEC. 928. DEFINITIONS.

14 *"In this title:*

15 "(1) ADVISORY COUNCIL.—The term 'Advisory
16 Council' means the Advisory Council on Healthcare
17 Research and Quality established under section 921.
18 "(2) AGENCY.—The term 'Agency' means the
19 Agency for Healthcare Research and Quality.
20 "(3) DIRECTOR.—The term 'Director' means the
21 Director for the Agency for Healthcare Research and

22 Quality.".

23 SEC. 403. REFERENCES.

24 Effective upon the date of enactment of this Act, any
25 reference in law to the "Agency for Health Care Policy and

1	(1) IN GENERAL.—Section $220(c)(1)(A)$ of the
2	Internal Revenue Code of 1986 (relating to eligible in-
3	dividual) is amended to read as follows:
4	"(A) IN GENERAL.—The term 'eligible indi-
5	vidual' means, with respect to any month, any
6	individual if—
7	"(i) such individual is covered under a
8	high deductible health plan as of the 1st day
9	of such month, and
10	"(ii) such individual is not, while cov-
11	ered under a high deductible health plan,
12	covered under any health plan—
13	"(I) which is not a high deduct-
14	ible health plan, and
15	"(II) which provides coverage for
16	any benefit which is covered under the
17	high deductible health plan.".
18	(2) Conforming Amendments.—
19	(A) Section $220(c)(1)$ of such Code is
20	amended by striking subparagraphs (C) and (D).
21	(B) Section 220(c) of such Code is amended
22	by striking paragraph (4) (defining small em-
23	ployer) and by redesignating paragraph (5) as
24	paragraph (4).

1	(C) Section 220(b) of such Code is amended
2	by striking paragraph (4) (relating to deduction
3	limited by compensation) and by redesignating
4	paragraphs (5), (6), and (7) as paragraphs (4),
5	(5), and (6), respectively.
6	(b) Removal of Limitation on Number of Tax-
7	PAYERS HAVING MEDICAL SAVINGS ACCOUNTS.—
8	(1) IN GENERAL.—Section 220 of the Internal
9	Revenue Code of 1986 (relating to medical savings ac-
10	counts) is amended by striking subsections (i) and (j).
11	(2) Medicare+choice.—Section 138 of such
12	Code (relating to Medicare+Choice MSA) is amended
13	by striking subsection (f).
14	(c) Reduction in High Deductible Plan Minimum
15	Annual Deductible.—
16	(1) IN GENERAL.—Subparagraph (A) of section
17	220(c)(2) of such Code (defining high deductible
18	health plan) is amended—
19	(A) by striking "\$1,500" and inserting
20	"\$1,000", and
21	(B) by striking "\$3,000" in clause (ii) and
22	inserting ''\$2,000''.
23	(2) Conforming Amendment.—Subsection (g)
24	of section 220 of such Code is amended—

1	(A) by striking "1998" and inserting
2	"1999"; and
3	(B) by striking "1997" and inserting
4	"1998".
5	(d) Increase in Contribution Limit to 100 Per-
6	CENT OF ANNUAL DEDUCTIBLE.—
7	(1) IN GENERAL.—Section 220(b)(2) of the Inter-
8	nal Revenue Code of 1986 (relating to monthly limi-
9	tation) is amended to read as follows:
10	"(2) MONTHLY LIMITATION.—The monthly limi-
11	tation for any month is the amount equal to $\frac{1}{12}$ of
12	the annual deductible of the high deductible health
13	plan of the individual.".
14	(2) Conforming Amendment.—Section
15	220(d)(1)(A) of such Code is amended by striking "75
16	percent of".
17	(e) Limitation on Additional Tax on Distribu-
18	tions Not Used for Qualified Medical Expenses.—
19	Section 220(f)(4) of the Internal Revenue Code of 1986 (re-
20	lating to additional tax on distributions not used for quali-
21	fied medical expenses) is amended by adding at the end the
22	following:
23	"(D) Exception in case of sufficient
24	Account balance.—Subparagraph (A) shall

25 not apply to any payment or distribution in

1	any taxable year, but only to the extent such
2	payment or distribution does not reduce the fair
3	market value of the assets of the medical savings
4	account to an amount less than the annual de-
5	ductible for the high deductible health plan of the
6	account holder (determined as of January 1 of
7	the calendar year in which the taxable year be-
8	gins).".
9	(f) TREATMENT OF NETWORK-BASED MANAGED CARE
10	PLANS.—Section 220(c)(2)(B) of the Internal Revenue Code
11	of 1986 (relating to special rules for high deductible health
12	plans) is amended by adding at the end the following:
13	"(iii) TREATMENT OF NETWORK-BASED
14	MANAGED CARE PLANS.—A plan that pro-
15	vides health care services through a network
16	of contracted or affiliated health care pro-
17	viders, if the benefits provided when services
18	are obtained through network providers
19	meet the requirements of subparagraph (A) ,
20	shall not fail to be treated as a high deduct-
21	ible health plan by reason of providing ben-
22	efits for services rendered by providers who
23	are not members of the network, so long as
24	the annual deductible and annual limit on
25	out-of-pocket expenses applicable to services

1	received from non-network providers are not
2	lower than those applicable to services re-
3	ceived from the network providers.".
4	(g) EFFECTIVE DATE.—The amendments made by this
5	section shall apply to taxable years beginning after Decem-
6	ber 31, 1999.
7	SEC. 503. PERMITTING CONTRIBUTION TOWARDS MEDICAL
8	SAVINGS ACCOUNT THROUGH FEDERAL EM-
9	PLOYEES HEALTH BENEFITS PROGRAM
10	(FEHBP).
11	(a) Authority To Contract for Catastrophic
12	PLANS.—Section 8902 of title 5, United States Code, is
13	amended by adding at the end the following:
14	((p)(1) The Office shall contract under this chapter for
15	a catastrophic plan with any qualified carrier that—
16	"(A) offers such a plan; and
17	``(B) as of the date of enactment of the Patients'
18	Bill of Rights Plus Act, offers a health benefits plan
19	under this chapter.
20	"(2) The Office may contract under this chapter for
21	a catastrophic plan with any qualified carrier that—
22	"(A) offers such a plan; but
23	``(B) does not satisfy the requirement under
24	paragraph (1)(B).".

(b) GOVERNMENT CONTRIBUTION TO MEDICAL SAV 2 INGS ACCOUNT.—

3 (1) IN GENERAL.—Section 8906 of title 5,
4 United States Code, is amended by adding at the end
5 the following:

6 "(j)(1) In the case of an employee or annuitant who 7 is enrolled in a catastrophic plan described by section 8 8903(5), there shall be a Government contribution under 9 this subsection to a medical savings account established or 10 maintained for the benefit of the individual. The contribu-11 tion under this subsection shall be in addition to the Gov-12 ernment contribution under subsection (b).

13 "(2) The amount of the Government contribution
14 under this subsection with respect to an individual is equal
15 to the amount by which—

16 "(A) the maximum contribution allowed under
17 subsection (b)(1) with respect to any employee or an18 nuitant, exceeds

"(B) the amount of the Government contribution
actually made with respect to the individual under
subsection (b) for coverage under the catastrophic
plan.

23 "(3) The Government contributions under this sub24 section shall be paid into a medical savings account (des25 ignated by the individual involved) in a manner that is

specified by the Office and consistent with the timing of
 contributions under subsection (b).

3 "(4) Subsections (f) and (g) shall apply to contribu4 tions under this section in the same manner as they apply
5 to contributions under subsection (b).

6 "(5) For the purpose of this subsection, the term 'med7 ical savings account' has the meaning given such term by
8 section 220(d) of the Internal Revenue Code of 1986.".

9 (2) ALLOWING PAYMENT OF FULL AMOUNT OF 10 CHARGE FOR CATASTROPHIC PLAN.—Section 11 8906(b)(2) of such title is amended by inserting "(or 12 100 percent of the subscription charge in the case of 13 a catastrophic plan)" after "75 percent of the sub-14 scription charge".

15 (c) Offering of Catastrophic Plans.—

16 (1) IN GENERAL.—Section 8903 of title 5,
17 United States Code, is amended by adding at the end
18 the following:

"(5) CATASTROPHIC PLANS.—(A) One or more
plans described in paragraph (1), (2), or (3), but
which provide benefits of the types referred to by
paragraph (5) of section 8904(a), instead of the types
referred to in paragraphs (1), (2), and (3) of such section.

1	``(B) Nothing in this section shall be
2	considered—
3	"(i) to prevent a carrier from simulta-
4	neously offering a plan described by subpara-
5	graph (A) and a plan described by paragraph
6	(1) or (2);
7	"(ii) to require that a catastrophic plan
8	offer two levels of benefits; or
9	"(iii) to allow, in any contract year, for-
10	``(I) more than one plan to be offered
11	which satisfies both subparagraph (A) and
12	paragraph (1) (subject to clause (ii)); and
13	"(II) more than one plan which satis-
14	fies both subparagraph (A) and paragraph
15	(2) (subject to clause (ii)).".
16	(2) Types of benefits.—Section 8904(a) of
17	such title is amended by inserting after paragraph (4)
18	the following new paragraph:
19	"(5) CATASTROPHIC PLANS.—Benefits of the
20	types named under paragraph (1) or (2) of this sub-
21	section or both, except that the plan shall meet the an-
22	nual deductible and annual out-of-pocket expenses re-
23	quirements under section $220(c)(2)$ of the Internal
24	Revenue Code of 1986.".

1 (3) Determining level of government con-2 TRIBUTIONS.—Section 8906(b) of such title is amend-3 ed by adding at the end the following: "Subscription 4 charges for medical savings accounts shall be deemed to be the amount of Government contributions made 5 6 under subsection (j)(2).". 7 (d) Conforming Amendments.— 8 (1) Additional health benefits plans.— 9 Section 8903a of title 5, United States Code, is 10 amended by redesignating subsection (d) as subsection 11 (e) and by inserting after subsection (c) the following: 12 "(d) The plans under this section may include one or more plans, otherwise allowable under this section, that sat-13 isfy the requirements of clauses (i) and (ii) of section 14 15 8903(5)(A).".

16 (2) REFERENCE.—Section 8909(d) of title 5,
17 United States Code, is amended by striking
18 "8903a(d)" and inserting "8903a(e)".

(e) REFERENCES.—Section 8903 of title 5, United
States Code, is amended by adding at the end (as a flush
left sentence) the following:

22 "The Office shall prescribe regulations under which the re23 quirements of section 8902(c), 8902(n), 8909(e), and any
24 other provision of this chapter that applies with respect to
25 a plan described by paragraph (1), (2), (3), or (4) of this

section shall apply with respect to the corresponding plan
 under paragraph (5) of this section. Similar regulations
 shall be prescribed with respect to any plan under section
 4 8903a(d).".

5 (f) EFFECTIVE DATE.—The amendments made by this
6 section shall apply to contract terms beginning on or after
7 January 1, 2000.

8 SEC. 504. CARRYOVER OF UNUSED BENEFITS FROM CAFE-9 TERIA PLANS, FLEXIBLE SPENDING AR-10 RANGEMENTS, AND HEALTH FLEXIBLE 11 SPENDING ACCOUNTS.

(a) IN GENERAL.—Section 125 of the Internal Revenue
Code of 1986 (relating to cafeteria plans) is amended by
redesignating subsections (h) and (i) as subsections (i) and
(j) and by inserting after subsection (g) the following new
subsection:

17 "(h) Allowance of Carryovers of Unused Bene18 FITS TO LATER TAXABLE YEARS.—

19	"(1) IN GENERAL.—For purposes of this title—
20	"(A) notwithstanding subsection $(d)(2)$, a
21	plan or other arrangement shall not fail to be
22	treated as a cafeteria plan or flexible spending or
23	similar arrangement, and

1	``(B) no amount shall be required to be in-
2	cluded in gross income by reason of this section
3	or any other provision of this chapter,
4	solely because under such plan or other arrangement
5	any nontaxable benefit which is unused as of the close
6	of a taxable year may be carried forward to 1 or more
7	succeeding taxable years.
8	"(2) LIMITATION.—Paragraph (1) shall not
9	apply to amounts carried from a plan to the extent
10	such amounts exceed \$500 (applied on an annual
11	basis). For purposes of this paragraph, all plans and
12	arrangements maintained by an employer or any re-
13	lated person shall be treated as 1 plan.
14	"(3) Allowance of rollover.—
15	"(A) IN GENERAL.—In the case of any un-
16	used benefit described in paragraph (1) which
17	consists of amounts in a health flexible spending
18	account or dependent care flexible spending ac-
19	count, the plan or arrangement shall provide
20	that a participant may elect, in lieu of such car-
21	ryover, to have such amounts distributed to the
22	participant.
23	"(B) Amounts not included in in-
24	COME.—Any distribution under subparagraph
25	(A) shall not be included in gross income to the

1	extent that such amount is transferred in a
2	trustee-to-trustee transfer, or is contributed with-
3	in 60 days of the date of the distribution, to-
4	"(i) a qualified cash or deferred ar-
5	rangement described in section 401(k),
6	"(ii) a plan under which amounts are
7	contributed by an individual's employer for
8	an annuity contract described in section
9	403(b),
10	"(iii) an eligible deferred compensation
11	plan described in section 457, or
12	"(iv) a medical savings account (with-
13	in the meaning of section 220).
14	Any amount rolled over under this subparagraph
15	shall be treated as a rollover contribution for the
16	taxable year from which the unused amount
17	would otherwise be carried.
18	"(C) TREATMENT OF ROLLOVER.—Any
19	amount rolled over under subparagraph (B)
20	shall be treated as an eligible rollover under sec-
21	tion 220, 401(k), 403(b), or 457, whichever is ap-
22	plicable, and shall be taken into account in ap-
23	plying any limitation (or participation require-
24	ment) on employer or employee contributions

1	under such section or any other provision of this
2	chapter for the taxable year of the rollover.
3	"(4) Cost-of-living adjustment.—In the case
4	of any taxable year beginning in a calendar year
5	after 1999, the $$500$ amount under paragraph (2)
6	shall be adjusted at the same time and in the same
7	manner as under section $415(d)(2)$, except that the
8	base period taken into account shall be the calendar
9	quarter beginning October 1, 1998, and any increase
10	which is not a multiple of \$50 shall be rounded to the
11	next lowest multiple of \$50.
12	"(5) Applicability.—This subsection shall
13	apply to taxable years beginning after December 31,
14	1999.".
15	(b) EFFECTIVE DATE.—The amendments made by this
16	section shall apply to taxable years beginning after Decem-
17	ber 31, 1999.

TITLE VI—PROVISIONS RELAT- ING TO LONG-TERM CARE IN- SURANCE

4 SEC. 601. INCLUSION OF QUALIFIED LONG-TERM CARE IN5 SURANCE CONTRACTS IN CAFETERIA PLANS,
6 FLEXIBLE SPENDING ARRANGEMENTS, AND
7 HEALTH FLEXIBLE SPENDING ACCOUNTS.

8 (a) IN GENERAL.—Section 125(f) of the Internal Rev9 enue Code of 1986 (defining qualified benefits) is amended
10 by striking the last sentence and inserting the following:
11 "Such term includes any qualified long-term care insurance
12 contract.".

(b) EFFECTIVE DATE.—The amendment made by this
section shall apply to taxable years beginning after December 31, 1999.

16SEC. 602. DEDUCTION FOR PREMIUMS FOR LONG-TERM17CARE INSURANCE.

(a) IN GENERAL.—Part VII of subchapter B of chapter
1 of the Internal Revenue Code of 1986 (relating to additional itemized deductions) is amended by redesignating
section 222 as section 223 and by inserting after section
22 221 the following:

23 "SEC. 222. PREMIUMS FOR LONG-TERM CARE INSURANCE.

24 "(a) IN GENERAL.—In the case of an eligible indi-25 vidual, there shall be allowed as a deduction an amount equal to 100 percent of the amount paid during the taxable
 year for any coverage for qualified long-term care services
 (as defined in section 7702B(c)) or any qualified long-term
 care insurance contract (as defined in section 7702B(b))
 which constitutes medical care for the taxpayer, his spouse,
 and dependents.

7 "(b) LIMITATIONS.—

8 "(1) DEDUCTION NOT AVAILABLE TO INDIVID-9 UALS ELIGIBLE FOR EMPLOYER-SUBSIDIZED COV-10 ERAGE.—

11 "(A) IN GENERAL.—Except as provided in 12 subparagraph (B), subsection (a) shall not apply 13 to any taxpayer for any calendar month for 14 which the taxpayer is eligible to participate in 15 any plan which includes coverage for qualified 16 long-term care services (as so defined) or is a 17 qualified long-term care insurance contract (as 18 so defined) maintained by any employer (or 19 former employer) of the taxpayer or of the spouse 20 of the taxpayer.

21 "(B) CONTINUATION COVERAGE.—Coverage
22 shall not be treated as subsidized for purposes of
23 this paragraph if—

24 "(i) such coverage is continuation cov25 erage (within the meaning of section

14980B(f)) required to be provided by the2employer, and

3 "(ii) the taxpayer or the taxpayer's
4 spouse is required to pay a premium for
5 such coverage in an amount not less than
6 100 percent of the applicable premium
7 (within the meaning of section 4980B(f)(4))
8 for the period of such coverage.

9 "(2) LIMITATION ON LONG-TERM CARE PRE-10 MIUMS.—In the case of a qualified long-term care in-11 surance contract (as so defined), only eligible long-12 premiums (as defined term care in section 13 213(d)(10)) shall be taken into account under sub-14 section (a)(2).

15 "(c) SPECIAL RULES.—For purposes of this section—
16 "(1) COORDINATION WITH MEDICAL DEDUCTION,
17 ETC.—Any amount paid by a taxpayer for insurance
18 to which subsection (a) applies shall not be taken into
19 account in computing the amount allowable to the
20 taxpayer as a deduction under section 213(a).

21 "(2) DEDUCTION NOT ALLOWED FOR SELF-EM22 PLOYMENT TAX PURPOSES.—The deduction allowable
23 by reason of this section shall not be taken into ac24 count in determining an individual's net earnings
1	from self-employment (within the meaning of section
2	1402(a)) for purposes of chapter 2.".
3	(b) Conforming Amendments.—
4	(1) Subsection (a) of section 62 of the Internal
5	Revenue Code of 1986 is amended by inserting after
6	paragraph (17) the following:
7	"(18) Long-term care insurance costs of
8	CERTAIN INDIVIDUALS.—The deduction allowed by
9	section 222.".
10	(2) The table of sections for part VII of sub-
11	chapter B of chapter 1 of such Code is amended by
12	striking the last item and inserting the following:
	"Sec. 222. Premiums for long-term care insurance. "Sec. 223. Cross reference.".
13	(c) EFFECTIVE DATE.—The amendments made by this
14	section shall apply to taxable years beginning after Decem-
15	ber 31, 1999.
16	SEC. 603. STUDY OF LONG-TERM CARE NEEDS IN THE 21ST
17	CENTURY.
18	(a) IN GENERAL.—The Secretary of Health and
19	Human Services (referred to in this section as the "Sec-
20	retary") shall provide, in accordance with this section, for
21	a study in order to determine—
22	(1) future demand for long-term health care serv-
23	ices (including institutional and home and commu-

1	nity-based services) in the United States in order to
2	meet the needs in the 21st century; and
3	(2) long-term options to finance the provision of
4	such services.
5	(b) Details.—The study conducted under subsection
6	(a) shall include the following:
7	(1) An identification of the relevant demographic
8	characteristics affecting demand for long-term health
9	care services, at least through the year 2030.
10	(2) The viability and capacity of community-
11	based and other long-term health care services under
12	different federal programs, including through the
13	medicare and medicaid programs, grants to States,
14	housing services, and changes in tax policy.
15	(3) How to improve the quality of long-term
16	health care services.
17	(4) The integration of long-term health care serv-
18	ices for individuals between different classes of health
19	care providers (such as hospitals, nursing facilities,
20	and home care agencies) and different Federal pro-
21	grams (such as the medicare and medicaid pro-
22	grams).
23	(5) The possibility of expanding private sector
24	initiatives, including long-term care insurance, to
25	meet the need to finance such services.

1	(6) An examination of the effect of enactment of
2	the Health Insurance Portability and Accountability
3	Act of 1996 on the provision and financing of long-
4	term health care services, including on portability
5	and affordability of private long-term care insurance,
6	the impact of insurance options on low-income older
7	Americans, and the options for eligibility to improve
8	access to such insurance.
9	(7) The financial impact of the provision of
10	long-term health care services on caregivers and other
11	family members.
12	(c) Report and Recommendations.—
13	(1) IN GENERAL.—Not later than 1 year after
14	the date of the enactment of this Act, the Secretary
15	shall provide for a report on the study under this sec-
16	tion.
17	(2) Recommendations.—The report under
18	paragraph (1) shall include findings and rec-
19	ommendations regarding each of the following:
20	(A) The most effective and efficient manner
21	that the Federal government may use its re-
22	sources to educate the public on planning for
23	needs for long-term health care services.

1	(B) The public, private, and joint public-
2	private strategies for meeting identified needs for
3	long-term health care services.
4	(C) The role of States and local commu-
5	nities in the financing of long-term health care
6	services.
7	(3) Inclusion of cost estimates.—The report
8	under paragraph (1) shall include cost estimates of
9	the various options for which recommendations are
10	made.
11	(d) Conduct of Study.—
12	(1) Use of institute of medicine.—The Sec-
13	retary of Health and Human Services shall seek to
14	enter into an appropriate arrangement with the In-
15	stitute of Medicine of the National Academy of
16	Sciences to conduct the study under this section. If
17	such an arrangement cannot be made, the Secretary
18	may provide for the conduct of the study by any other
19	qualified non-governmental entity.
20	(2) CONSULTATION.—The study should be con-
21	ducted under this section in consultation with experts
22	from a wide-range of groups from the public and pri-
02	and a sector of

vate sectors.

	221
1	TITLE VII—INDIVIDUAL
2	RETIREMENT PLANS
3	SEC. 701. MODIFICATION OF INCOME LIMITS ON CONTRIBU-
4	TIONS AND ROLLOVERS TO ROTH IRAS.
5	(a) Increase in AGI Limit for Rollover Con-
6	TRIBUTIONS.—Clause (i) of section 408A(c)(3)(A) of the In-
7	ternal Revenue Code of 1986 (relating to rollover from
8	IRA), as redesignated by subsection (a), is amended by
9	striking "\$100,000" and inserting "\$1,000,000".
10	(b) Conforming Amendments.—
11	(1)(A) Subparagraph (B) of section $408A(c)(3)$
12	of the Internal Revenue Code of 1986, as redesignated
13	by subsection (a), is amended to read as follows:
14	"(B) Definition of adjusted gross in-
15	COME.—For purposes of subparagraph (A), ad-
16	justed gross income shall be determined—
17	"(i) after application of sections 86
18	and 469, and
19	"(ii) without regard to sections 135,
20	137, 221, and 911, the deduction allowable
21	under section 219, or any amount included
22	in gross income under subsection $(d)(3)$.".
23	(B) EFFECTIVE DATE.—The amendment made
24	by this paragraph shall apply to taxable years begin-
25	ning after December 31, 1999.

1	(2)(A) Subparagraph (B) of section $408A(c)(3)$
2	of such Code, as amended by paragraph (1), is
3	amended to read as follows:
4	"(B) DEFINITION OF ADJUSTED GROSS IN-
5	COME.—For purposes of subparagraph (A), ad-
6	justed gross income shall be determined—
7	"(i) after application of sections 86
8	and 469, and
9	"(ii) without regard to sections 135,
10	137, 221, and 911, the deduction allowable
11	under section 219, or any amount included
12	in gross income under subsection $(d)(3)$ or
13	by reason of a required distribution under
14	a provision described in paragraph (5).".
15	(B) EFFECTIVE DATE.—The amendment made
16	by this paragraph shall apply to taxable years begin-
17	ning after December 31, 2004.
18	(c) Effective Date.—Except as otherwise provided
19	in this section, the amendments made by this section shall
20	apply to taxable years beginning after December 31, 1999.

TITLE VIII—REVENUE PROVISIONS

223

1

2

3 SEC. 801. MODIFICATION TO FOREIGN TAX CREDIT 4 CARRYBACK AND CARRYOVER PERIODS.

5 (a) IN GENERAL.—Section 904(c) of the Internal Rev6 enue Code of 1986 (relating to limitation on credit) is
7 amended—

8 (1) by striking "in the second preceding taxable
9 year,", and

10 (2) by striking "or fifth" and inserting "fifth,
11 sixth, or seventh".

(b) EFFECTIVE DATE.—The amendment made by subsection (a) shall apply to credits arising in taxable years
beginning after December 31, 2001.

15 SEC. 802. LIMITATION ON USE OF NON-ACCRUAL EXPERI16 ENCE METHOD OF ACCOUNTING.

17 (a) IN GENERAL.—Section 448(d)(5) of the Internal
18 Revenue Code of 1986 (relating to special rule for services)
19 is amended—

20 (1) by inserting "in fields described in para21 graph (2)(A)" after "services by such person", and

(2) by inserting "CERTAIN PERSONAL" before
23 "SERVICES" in the heading.

24 (b) EFFECTIVE DATE.—

1	(1) IN GENERAL.—The amendments made by
2	this section shall apply to taxable years ending after
3	the date of the enactment of this Act.
4	(2) Change in method of accounting.—In
5	the case of any taxpayer required by the amendments
6	made by this section to change its method of account-
7	ing for its first taxable year ending after the date of
8	the enactment of this Act—
9	(A) such change shall be treated as initiated
10	by the taxpayer,
11	(B) such change shall be treated as made
12	with the consent of the Secretary of the Treasury,
13	and
14	(C) the net amount of the adjustments re-
15	quired to be taken into account by the taxpayer
16	under section 481 of the Internal Revenue Code
17	of 1986 shall be taken into account over a period
18	(not greater than 4 taxable years) beginning
19	with such first taxable year.
20	SEC. 803. RETURNS RELATING TO CANCELLATIONS OF IN-
21	DEBTEDNESS BY ORGANIZATIONS LENDING
22	MONEY.
23	(a) IN GENERAL.—Paragraph (2) of section $6050P(c)$
24	of the Internal Revenue Code of 1986 (relating to definitions
25	and special rules) is amended by striking "and" at the end

of subparagraph (B), by striking the period at the end of 1 subparagraph (C) and inserting ", and", and by inserting 2 after subparagraph (C) the following new subparagraph: 3 4 (D) any organization a significant trade 5 or business of which is the lending of money.". 6 (b) EFFECTIVE DATE.—The amendment made by sub-7 section (a) shall apply to discharges of indebtedness after 8 December 31, 1999. 9 SEC. 804. EXTENSION OF INTERNAL REVENUE SERVICE 10 USER FEES.

(a) IN GENERAL.—Chapter 77 of the Internal Revenue
 Code of 1986 (relating to miscellaneous provisions) is
 amended by adding at the end the following new section:
 "SEC. 7527. INTERNAL REVENUE SERVICE USER FEES.

15 "(a) GENERAL RULE.—The Secretary shall establish
16 a program requiring the payment of user fees for—

17 "(1) requests to the Internal Revenue Service for
18 ruling letters, opinion letters, and determination let19 ters, and

20 "(2) other similar requests.

21 "(b) PROGRAM CRITERIA.—

22 "(1) IN GENERAL.—The fees charged under the
23 program required by subsection (a)—

24 "(A) shall vary according to categories (or
25 subcategories) established by the Secretary,

1	((B) shall be determined after taking into
2	account the average time for (and difficulty of)
3	complying with requests in each category (and
4	subcategory), and
5	"(C) shall be payable in advance.
6	"(2) EXEMPTIONS, ETC.—The Secretary shall
7	provide for such exemptions (and reduced fees) under
8	such program as the Secretary determines to be ap-
9	propriate.
10	"(3) Average fee requirement.—The average
11	fee charged under the program required by subsection
10	(a) shall not be less than the amount determined
12	(a) shall not be less than the amount determined
12 13	(a) shall not be less than the amount actermined under the following table:
	under the following table:"CategoryAverage FeeEmployee plan ruling and opinion
13	under the following table:"CategoryAverage FeeEmployee plan ruling and opinion\$250Exempt organization ruling\$350Employee plan determination\$300Exempt organization determination\$275Chief counsel ruling\$200.
13 14	under the following table:"CategoryAverage FeeEmployee plan ruling and opinion\$250Exempt organization ruling\$350Employee plan determination\$300Exempt organization determination\$275Chief counsel ruling\$200."(c) TERMINATION.—No fee shall be imposed under
13 14 15	under the following table:"CategoryAverage FeeEmployee plan ruling and opinion\$250Exempt organization ruling\$350Employee plan determination\$300Exempt organization determination\$275Chief counsel ruling\$200."(c) TERMINATION.—No fee shall be imposed underthis section with respect to requests made after September
13 14 15 16	under the following table:"CategoryAverage FeeEmployee plan ruling and opinion\$250Exempt organization ruling\$350Employee plan determination\$300Exempt organization determination\$275Chief counsel ruling\$200."(c) TERMINATION.—No fee shall be imposed underthis section with respect to requests made after September30, 2009.".
13 14 15 16 17	under the following table: "Category Average Fee Employee plan ruling and opinion
 13 14 15 16 17 18 	under the following table:"CategoryAverage FeeEmployee plan ruling and opinion\$250Exempt organization ruling\$350Employee plan determination\$300Exempt organization determination\$275Chief counsel ruling\$200."(c) TERMINATION.—No fee shall be imposed underthis section with respect to requests made after September30, 2009.".(b) CONFORMING AMENDMENTS.—(1) The table of sections for chapter 77 of the In-

1 (2) Section 10511 of the Revenue Act of 1987 is 2 repealed. 3 (c) EFFECTIVE DATE.—The amendments made by this 4 section shall apply to requests made after the date of the enactment of this Act. 5 6 SEC. 805. PROPERTY SUBJECT TO A LIABILITY TREATED IN 7 SAME MANNER AS ASSUMPTION OF LIABIL-8 ITY. 9 (a) Repeal of Property Subject to a Liability 10 TEST.— 11 (1) SECTION 357.—Section 357(a)(2) of the Inter-12 nal Revenue Code of 1986 (relating to assumption of liability) is amended by striking ", or acquires from 13 14 the taxpayer property subject to a liability". 15 (2) SECTION 358.—Section 358(d)(1) of such 16 Code (relating to assumption of liability) is amended 17 by striking "or acquired from the taxpayer property 18 subject to a liability". 19 (3) Section 368.— 20 (A) Section 368(a)(1)(C) of such Code is 21 amended by striking ", or the fact that property 22 acquired is subject to a liability,". 23 (B)The last sentence ofsection 24 368(a)(2)(B) of such Code is amended by strik-25 ing ", and the amount of any liability to which

1	any property acquired from the acquiring cor-
2	poration is subject,".
3	(b) Clarification of Assumption of Liability.—
4	(1) In general.—Section 357 of the Internal
5	Revenue Code of 1986 is amended by adding at the
6	end the following new subsection:
7	"(d) Determination of Amount of Liability As-
8	SUMED.—
9	"(1) IN GENERAL.—For purposes of this section,
10	section $358(d)$, section $362(d)$, section $368(a)(1)(C)$,
11	and section $368(a)(2)(B)$, except as provided in
12	regulations—
13	"(A) a recourse liability (or portion thereof)
14	shall be treated as having been assumed if, as de-
15	termined on the basis of all facts and cir-
16	cumstances, the transferee has agreed to, and is
17	expected to, satisfy such liability (or portion),
18	whether or not the transferor has been relieved of
19	such liability, and
20	"(B) except to the extent provided in para-
21	graph (2), a nonrecourse liability shall be treated
22	as having been assumed by the transferee of any
23	asset subject to such liability.
24	"(2) Exception for nonrecourse liabil-
25	ITY.—The amount of the nonrecourse liability treated

1	as described in paragraph $(1)(B)$ shall be reduced by
2	the lesser of—
3	((A) the amount of such liability which an
4	owner of other assets not transferred to the trans-
5	feree and also subject to such liability has agreed
6	with the transferee to, and is expected to, satisfy,
7	OT
8	"(B) the fair market value of such other as-
9	sets (determined without regard to section
10	7701(g)).
11	"(3) REGULATIONS.—The Secretary shall pre-
12	scribe such regulations as may be necessary to carry
13	out the purposes of this subsection and section $362(d)$.
14	The Secretary may also prescribe regulations which
15	provide that the manner in which a liability is treat-
16	ed as assumed under this subsection is applied, where
17	appropriate, elsewhere in this title.".
18	(2) LIMITATION ON BASIS INCREASE ATTRIB-
19	UTABLE TO ASSUMPTION OF LIABILITY.—Section 362
20	of such Code is amended by adding at the end the fol-
21	lowing new subsection:
22	"(d) Limitation on Basis Increase Attributable
23	to Assumption of Liability.—
24	"(1) IN GENERAL.—In no event shall the basis of
25	any property be increased under subsection (a) or (b)

1	above the fair market value of such property (deter-
2	mined without regard to section $7701(g)$) by reason of
3	any gain recognized to the transferor as a result of
4	the assumption of a liability.
5	"(2) TREATMENT OF GAIN NOT SUBJECT TO
6	TAX.—Except as provided in regulations, if—
7	"(A) gain is recognized to the transferor as
8	a result of an assumption of a nonrecourse li-
9	ability by a transferee which is also secured by
10	assets not transferred to such transferee, and
11	"(B) no person is subject to tax under this
12	title on such gain,
13	then, for purposes of determining basis under sub-
14	sections (a) and (b), the amount of gain recognized by
15	the transferor as a result of the assumption of the li-
16	ability shall be determined as if the liability assumed
17	by the transferee equaled such transferee's ratable por-
18	tion of such liability determined on the basis of the
19	relative fair market values (determined without re-
20	gard to section $7701(g)$) of all of the assets subject to
21	such liability.".
22	(c) Application to Provisions Other Than Sub-
23	Chapter C.—
24	(1) Section 584.—Section 584(h)(3) of the Inter-
25	nal Revenue Code of 1986 is amended—

1	(A) by striking ", and the fact that any
2	property transferred by the common trust fund is
3	subject to a liability," in subparagraph (A), and
4	(B) by striking clause (ii) of subparagraph
5	(B) and inserting:
6	"(ii) Assumed liabilities.—For pur-
7	poses of clause (i), the term 'assumed liabil-
8	ities' means any liability of the common
9	trust fund assumed by any regulated invest-
10	ment company in connection with the
11	transfer referred to in paragraph $(1)(A)$.
12	"(C) Assumption.—For purposes of this
13	paragraph, in determining the amount of any li-
14	ability assumed, the rules of section 357(d) shall
15	apply.".
16	(2) Section 1031.—The last sentence of section
17	1031(d) of such Code is amended—
18	(A) by striking "assumed a liability of the
19	taxpayer or acquired from the taxpayer property
20	subject to a liability" and inserting "assumed
21	(as determined under section 357(d)) a liability
22	of the taxpayer", and
23	(B) by striking "or acquisition (in the
24	amount of the liability)".
25	(d) Conforming Amendments.—

1	(1) Section $351(h)(1)$ of the Internal Revenue
2	Code of 1986 is amended by striking ", or acquires
3	property subject to a liability,".
4	(2) Section 357 of such Code is amended by
5	striking "or acquisition" each place it appears in
6	subsection (a) or (b).
7	(3) Section 357(b)(1) of such Code is amended by
8	striking "or acquired".
9	(4) Section $357(c)(1)$ of such Code is amended by
10	striking ", plus the amount of the liabilities to which
11	the property is subject,".
12	(5) Section $357(c)(3)$ of such Code is amended by
13	striking "or to which the property transferred is sub-
14	ject".
15	(6) Section $358(d)(1)$ of such Code is amended
16	by striking "or acquisition (in the amount of the li-
17	ability)".
18	(e) EFFECTIVE DATE.—The amendments made by this
19	section shall apply to transfers after October 19, 1998.
20	SEC. 806. CHARITABLE SPLIT-DOLLAR LIFE INSURANCE, AN-
21	NUITY, AND ENDOWMENT CONTRACTS.
22	(a) IN GENERAL.—Subsection (f) of section 170 of the
23	Internal Revenue Code of 1986 (relating to disallowance of
24	deduction in certain cases and special rules) is amended
25	by adding at the end the following new paragraph:

1	"(10) Split-dollar life insurance, annuity,
2	AND ENDOWMENT CONTRACTS.—
3	"(A) IN GENERAL.—Nothing in this section
4	or in section 545(b)(2), 556(b)(2), 642(c), 2055,
5	2106(a)(2), or 2522 shall be construed to allow
6	a deduction, and no deduction shall be allowed,
7	for any transfer to or for the use of an organiza-
8	tion described in subsection (c) if in connection
9	with such transfer—
10	"(i) the organization directly or indi-
11	rectly pays, or has previously paid, any
12	premium on any personal benefit contract
13	with respect to the transferor, or
14	"(ii) there is an understanding or ex-
15	pectation that any person will directly or
16	indirectly pay any premium on any per-
17	sonal benefit contract with respect to the
18	transferor.
19	"(B) PERSONAL BENEFIT CONTRACT.—For
20	purposes of subparagraph (A), the term 'personal
21	benefit contract' means, with respect to the
22	transferor, any life insurance, annuity, or en-
23	dowment contract if any direct or indirect bene-

25 member of the transferor's family, or any other

ficiary under such contract is the transferor, any

24

1	person (other than an organization described in
2	subsection (c)) designated by the transferor.
3	"(C) Application to charitable remain-
4	DER TRUSTS.—In the case of a transfer to a
5	trust referred to in subparagraph (E) , references
6	in subparagraphs (A) and (F) to an organiza-
7	tion described in subsection (c) shall be treated
8	as a reference to such trust.
9	"(D) Exception for certain annuity
10	CONTRACTS.—If, in connection with a transfer to
11	or for the use of an organization described in
12	subsection (c), such organization incurs an obli-
13	gation to pay a charitable gift annuity (as de-
14	fined in section $501(m)$) and such organization
15	purchases any annuity contract to fund such ob-
16	ligation, persons receiving payments under the
17	charitable gift annuity shall not be treated for
18	purposes of subparagraph (B) as indirect bene-
19	ficiaries under such contract if—
20	((i) such organization possesses all of
21	the incidents of ownership under such con-
22	tract,
23	((ii) such organization is entitled to
24	all the payments under such contract, and

1	"(iii) the timing and amount of pay-
2	ments under such contract are substantially
3	the same as the timing and amount of pay-
4	ments to each such person under such obli-
5	gation (as such obligation is in effect at the
6	time of such transfer).
7	"(E) Exception for certain contracts
8	HELD BY CHARITABLE REMAINDER TRUSTS.—A
9	person shall not be treated for purposes of sub-
10	paragraph (B) as an indirect beneficiary under
11	any life insurance, annuity, or endowment con-
12	tract held by a charitable remainder annuity
13	trust or a charitable remainder unitrust (as de-
14	fined in section $664(d)$) solely by reason of being
15	entitled to any payment referred to in para-
16	graph (1)(A) or (2)(A) of section 664(d) if—
17	"(i) such trust possesses all of the inci-
18	dents of ownership under such contract, and
19	"(ii) such trust is entitled to all the
20	payments under such contract.
21	"(F) Excise tax on premiums paid.—
22	"(i) IN GENERAL.—There is hereby im-
23	posed on any organization described in sub-
24	section (c) an excise tax equal to the pre-
25	miums paid by such organization on any

1	life insurance, annuity, or endowment con-
2	tract if the payment of premiums on such
3	contract is in connection with a transfer for
4	which a deduction is not allowable under
5	subparagraph (A), determined without re-
6	gard to when such transfer is made.
7	"(ii) PAYMENTS BY OTHER PER-
8	sons.—For purposes of clause (i), pay-
9	ments made by any other person pursuant
10	to an understanding or expectation referred
11	to in subparagraph (A) shall be treated as
12	made by the organization.
13	"(iii) Reporting.—Any organization
14	on which tax is imposed by clause (i) with
15	respect to any premium shall file an annual
16	return which includes—
17	((I) the amount of such premiums
18	paid during the year and the name
19	and TIN of each beneficiary under the
20	contract to which the premium relates,
21	and
22	``(II) such other information as
23	the Secretary may require.
24	The penalties applicable to returns required
25	under section 6033 shall apply to returns

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1	required under this clause. Returns required
2	under this clause shall be furnished at such
3	time and in such manner as the Secretary
4	shall by forms or regulations require.
5	"(iv) Certain rules to Apply.—The
6	tax imposed by this subparagraph shall be
7	treated as imposed by chapter 42 for pur-
8	poses of this title other than subchapter B
9	of chapter 42.
10	"(G) Special rule where state re-
11	QUIRES SPECIFICATION OF CHARITABLE GIFT AN-
12	NUITANT IN CONTRACT.—In the case of an obli-
13	gation to pay a charitable gift annuity referred
14	to in subparagraph (D) which is entered into
15	under the laws of a State which requires, in
16	order for the charitable gift annuity to be exempt
17	from insurance regulation by such State, that
18	each beneficiary under the charitable gift annu-
19	ity be named as a beneficiary under an annuity
20	contract issued by an insurance company au-
21	thorized to transact business in such State, the
22	requirements of clauses (i) and (ii) of subpara-
23	graph (D) shall be treated as met if—
24	"(i) such State law requirement was in
25	effect on February 8, 1999,

((/::) and here the second sec
"(ii) each such beneficiary under the
charitable gift annuity is a bona fide resi-
dent of such State at the time the obligation
to pay a charitable gift annuity is entered
into, and
"(iii) the only persons entitled to pay-
ments under such contract are persons enti-
tled to payments as beneficiaries under such
obligation on the date such obligation is en-
tered into.
"(H) REGULATIONS.—The Secretary shall
prescribe such regulations as may be necessary
or appropriate to carry out the purposes of this
paragraph, including regulations to prevent the
avoidance of such purposes.".
(b) Effective Date.—
(1) IN GENERAL.—Except as otherwise provided
in this section, the amendment made by this section
shall apply to transfers made after February 8, 1999.
(2) Excise tax.—Except as provided in para-
graph (3) of this subsection, section $170(f)(10)(F)$ of
the Internal Revenue Code of 1986 (as added by this
section) shall apply to premiums paid after the date
of the enactment of this Act.

1	(3) REPORTING.—Clause (iii) of such section
2	170(f)(10)(F) shall apply to premiums paid after
3	February 8, 1999 (determined as if the tax imposed
4	by such section applies to premiums paid after such
5	date).
6	SEC. 807. TRANSFER OF EXCESS DEFINED BENEFIT PLAN
7	ASSETS FOR RETIREE HEALTH BENEFITS.
8	(a) EXTENSION.—
9	(1) IN GENERAL.—Section 420(b)(5) of the Inter-
10	nal Revenue Code of 1986 (relating to expiration) is
11	amended by striking "in any taxable year beginning
12	after December 31, 2000" and inserting "made after
13	September 30, 2009".
14	(2) Conforming Amendments.—
15	(A) Section $101(e)(3)$ of the Employee Re-
16	tirement Income Security Act of 1974 (29 U.S.C.
17	1021(e)(3)) is amended by striking "1995" and
18	inserting "2001".
19	(B) Section $403(c)(1)$ of such Act (29 U.S.C.
20	1103(c)(1)) is amended by striking "1995" and in-
21	serting "2001".
22	(C) Paragraph (13) of section 408(b) of such Act
23	(29 U.S.C. 1108(b)(13)) is amended—

1	/·\
1	(i) by striking "in a taxable year beginning
2	before January 1, 2001" and inserting "made
3	before October 1, 2009", and
4	(ii) by striking "1995" and inserting
5	"2001".
6	(b) Application of Minimum Cost Require-
7	MENTS.—
8	(1) IN GENERAL.—Section 420(c)(3) of the Inter-
9	nal Revenue Code of 1986 is amended to read as fol-
10	lows:
11	"(3) Minimum cost requirements.—
12	"(A) IN GENERAL.—The requirements of
13	this paragraph are met if each group health plan
14	or arrangement under which applicable health
15	benefits are provided provides that the applicable
16	employer cost for each taxable year during the
17	cost maintenance period shall not be less than
18	the higher of the applicable employer costs for
19	each of the 2 taxable years immediately pre-
20	ceding the taxable year of the qualified transfer.
21	"(B) Applicable employer cost.—For
22	purposes of this paragraph, the term 'applicable
23	employer cost' means, with respect to any tax-
24	able year, the amount determined by dividing—

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1	"(i) the qualified current retiree health
2	liabilities of the employer for such taxable
3	year determined—
4	"(I) without regard to any reduc-
5	tion under subsection $(e)(1)(B)$, and
6	"(II) in the case of a taxable year
7	in which there was no qualified trans-
8	fer, in the same manner as if there had
9	been such a transfer at the end of the
10	taxable year, by
11	"(ii) the number of individuals to
12	whom coverage for applicable health benefits
13	was provided during such taxable year.
14	"(C) Election to compute cost sepa-
15	RATELY.—An employer may elect to have this
16	paragraph applied separately with respect to in-
17	dividuals eligible for benefits under title XVIII of
18	the Social Security Act at any time during the
19	taxable year and with respect to individuals not
20	so eligible.
21	"(D) Cost maintenance period.—For
22	purposes of this paragraph, the term 'cost main-
23	tenance period' means the period of 5 taxable
24	years beginning with the taxable year in which
25	the qualified transfer occurs. If a taxable year is

1	in 2 or more overlapping cost maintenance peri-
2	ods, this paragraph shall be applied by taking
3	into account the highest applicable employer cost
4	required to be provided under subparagraph (A)
5	for such taxable year.".
6	(2) Conforming Amendments.—
7	(A) Section $420(b)(1)(C)(iii)$ of such Code is
8	amended by striking ''benefits'' and inserting
9	"cost".
10	(B) Section $420(e)(1)(D)$ of such Code is
11	amended by striking "and shall not be subject to
12	the minimum benefit requirements of subsection
13	(c)(3)" and inserting "or in calculating applica-
14	ble employer cost under subsection $(c)(3)(B)$ ".
15	(c) EFFECTIVE DATE.—The amendments made by this
16	section shall apply to qualified transfers occurring after De-
17	cember 31, 2000, and before October 1, 2009.
18	SEC. 808. LIMITATIONS ON WELFARE BENEFIT FUNDS OF 10
19	OR MORE EMPLOYER PLANS.
20	(a) Benefits to Which Exception Applies.—Sec-
21	tion 419A(f)(6)(A) of the Internal Revenue Code of 1986
22	(relating to exception for 10 or more employer plans) is
23	amended to read as follows:
24	"(A) IN GENERAL.—This subpart shall not
25	apply to a welfare benefit fund which is part of

1	a 10 or more employer plan if the only benefits
2	provided through the fund are 1 or more of the
3	following:
4	"(i) Medical benefits.
5	"(ii) Disability benefits.
6	"(iii) Group term life insurance bene-
7	fits which do not provide for any cash sur-
8	render value or other money that can be
9	paid, assigned, borrowed, or pledged for col-
10	lateral for a loan.
11	The preceding sentence shall not apply to any
12	plan which maintains experience-rating arrange-
13	ments with respect to individual employers.".
14	(b) Limitation on Use of Amounts for Other
15	Purposes.—Section 4976(b) of the Internal Revenue Code
16	of 1986 (defining disqualified benefit) is amended by add-
17	ing at the end the following new paragraph:
18	"(5) Special rule for 10 or more employer
19	PLANS EXEMPTED FROM PREFUNDING LIMITS.—For
20	purposes of paragraph (1)(C), if—
21	((A) subpart D of part I of subchapter D
22	of chapter 1 does not apply by reason of section
23	419A(f)(6) to contributions to provide 1 or more
24	welfare benefits through a welfare benefit fund
25	under a 10 or more employer plan, and

1	``(B) any portion of the welfare benefit fund
2	attributable to such contributions is used for a
3	purpose other than that for which the contribu-
4	tions were made,
5	then such portion shall be treated as reverting to the
6	benefit of the employers maintaining the fund.".
7	(c) EFFECTIVE DATE.—The amendments made by this
8	section shall apply to contributions paid or accrued after
9	the date of the enactment of this Act, in taxable years end-
10	ing after such date.
11	SEC. 809. MODIFICATION OF INSTALLMENT METHOD AND
12	REPEAL OF INSTALLMENT METHOD FOR AC-
13	CRUAL METHOD TAXPAYERS.
14	(a) Repeal of Installment Method for Accrual
14 15	(a) Repeal of Installment Method for Accrual Basis Taxpayers.—
15	BASIS TAXPAYERS.—
15 16	BASIS TAXPAYERS.— (1) IN GENERAL.—Subsection (a) of section 453
15 16 17	BASIS TAXPAYERS.— (1) IN GENERAL.—Subsection (a) of section 453 of the Internal Revenue Code of 1986 (relating to in-
15 16 17 18	BASIS TAXPAYERS.— (1) IN GENERAL.—Subsection (a) of section 453 of the Internal Revenue Code of 1986 (relating to in- stallment method) is amended to read as follows:
15 16 17 18 19	BASIS TAXPAYERS.— (1) IN GENERAL.—Subsection (a) of section 453 of the Internal Revenue Code of 1986 (relating to in- stallment method) is amended to read as follows: "(a) USE OF INSTALLMENT METHOD.—
15 16 17 18 19 20	BASIS TAXPAYERS.— (1) IN GENERAL.—Subsection (a) of section 453 of the Internal Revenue Code of 1986 (relating to in- stallment method) is amended to read as follows: "(a) USE OF INSTALLMENT METHOD.— "(1) IN GENERAL.—Except as otherwise provided
15 16 17 18 19 20 21	BASIS TAXPAYERS.— (1) IN GENERAL.—Subsection (a) of section 453 of the Internal Revenue Code of 1986 (relating to in- stallment method) is amended to read as follows: "(a) USE OF INSTALLMENT METHOD.— "(1) IN GENERAL.—Except as otherwise provided in this section, income from an installment sale shall
 15 16 17 18 19 20 21 22 	BASIS TAXPAYERS.— (1) IN GENERAL.—Subsection (a) of section 453 of the Internal Revenue Code of 1986 (relating to in- stallment method) is amended to read as follows: "(a) USE OF INSTALLMENT METHOD.— "(1) IN GENERAL.—Except as otherwise provided in this section, income from an installment sale shall be taken into account for purposes of this title under

1	stallment sale if such income would be reported under
2	an accrual method of accounting without regard to
3	this section. The preceding sentence shall not apply to
4	a disposition described in subparagraph (A) or (B) of
5	subsection $(l)(2)$.".

6 (2) CONFORMING AMENDMENTS.—Sections
7 453(d)(1), 453(i)(1), and 453(k) of such Code are each
8 amended by striking "(a)" each place it appears and
9 inserting "(a)(1)".

(b) MODIFICATION OF PLEDGE RULES.—Paragraph 10 11 (4) of section 453A(d) of the Internal Revenue Code of 1986 12 (relating to pledges, etc., of installment obligations) is amended by adding at the end the following: "A payment 13 14 shall be treated as directly secured by an interest in an 15 installment obligation to the extent an arrangement allows the taxpayer to satisfy all or a portion of the indebtedness 16 17 with the installment obligation.".

(c) EFFECTIVE DATE.—The amendments made by this
section shall apply to sales or other dispositions occurring
on or after the date of the enactment of this Act.

1	SEC. 810. INCLUSION OF CERTAIN VACCINES AGAINST
2	STREPTOCOCCUS PNEUMONIAE TO LIST OF
3	TAXABLE VACCINES.
4	(a) IN GENERAL.—Section 4132(a)(1) of the Internal
5	Revenue Code of 1986 (defining taxable vaccine) is amended
6	by adding at the end the following new subparagraph:
7	"(L) Any conjugate vaccine against strepto-
8	coccus pneumoniae.".
9	(b) Effective Date.—
10	(1) SALES.—The amendment made by this sec-
11	tion shall apply to vaccine sales beginning on the day
12	after the date on which the Centers for Disease Con-
13	trol makes a final recommendation for routine ad-
14	ministration to children of any conjugate vaccine
15	against streptococcus pneumoniae.
16	(2) DELIVERIES.—For purposes of paragraph
17	(1), in the case of sales on or before the date described
18	in such paragraph for which delivery is made after
19	such date, the delivery date shall be considered the
20	sale date.

TITLE IX—MISCELLANEOUS PROVISIONS

1

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3 SEC. 901. MEDICARE COMPETITIVE PRICING DEMONSTRA-4 TION PROJECT.

5 (a) FINDING.—The Senate finds that implementing
6 competitive pricing in the medicare program under title
7 XVIII of the Social Security Act is an important goal.

8 (b) PROHIBITION ON IMPLEMENTATION OF PROJECT IN CERTAIN AREAS.—Notwithstanding subsection (b) of sec-9 10 tion 4011 of the Balanced Budget Act of 1997 (Public Law 11 105–33)), the Secretary of Health and Human Services 12 may not implement the Medicare Competitive Pricing Demonstration Project (operated by the Secretary of Health and 13 14 Human Services pursuant to such section) in Kansas City, Missouri or Kansas City, Kansas, or in any area in Ari-15 16 zona.

(c) MORATORIUM ON IMPLEMENTATION OF PROJECT IN
ANY AREA UNTIL JANUARY, 1, 2001.—Notwithstanding any
provision of section 4011 of the Balanced Budget Act of
1997 (Public Law 105–33)), the Secretary of Health and
Human Services may not implement the Medicare Competitive Pricing Demonstration Project in any area before January 1, 2001.

24 (d) Study and Report to Congress.—

1	(1) STUDY.—The Secretary of Health and
2	Human Services, in conjunction with the Competitive
3	Pricing Advisory Committee, shall conduct a study on
4	the different approaches of implementing the Medicare
5	Competitive Pricing Demonstration Project on a vol-
6	untary basis.
7	(2) REPORT.—Not later than June 30, 2000, the
8	Secretary of Health and Human Services shall sub-
9	mit a report to Congress which shall contain a de-
10	tailed description of the study conducted under para-
11	graph (1), together with the recommendations of the
12	Secretary and the Competitive Pricing Advisory Com-
13	mittee regarding the implementation of the Medicare
14	Competitive Pricing Demonstration Project.
	Attest:

Secretary.



AMENDMENT

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- HR 2990 EAS—12
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