§ 54.9802–3T Additional requirements prohibiting discrimination based on genetic information (temporary).

(a) Definitions. Unless otherwise provided, the definitions in this paragraph (a) govern in applying the provisions of this section.

(1) Collect means, with respect to information, to request, require, or purchase such information.

(2) Family member means, with respect to an individual —

(i) A dependent (as defined for purposes of §54.9801–2) of the individual; or

(ii) Any other person who is a first-degree, second-degree, third-degree, or fourth-degree relative of the individual or of a dependent of the individual. Relatives by affinity (such as by marriage or adoption) are treated the same as relatives by consanguinity (that is, relatives who share a common biological ancestor). In determining the degree of the relationship, relatives by less than full consanguinity (such as half-siblings, who share only one parent) are treated the same as relatives by full consanguinity (such as siblings who share both parents).

(A) First-degree relatives include parents, spouses, siblings, and children.

(B) Second-degree relatives include grandparents, grandchildren, aunts, uncles, nephews, and nieces.

(C) Third-degree relatives include great-grandparents, great-grandchildren, great aunts, great uncles, and first cousins.

(D) Fourth-degree relatives include great-great grandparents, great-great grandchildren, and children of first cousins.

(3) Genetic information means—

(i) Subject to paragraphs (a)(3)(ii) and (a)(3)(iii) of this section, with respect to an individual, information about—

(A) The individual’s genetic tests (as defined in paragraph (a)(5) of this section);

(B) The genetic tests of family members of the individual;

(C) The manifestation (as defined in paragraph (a)(6) of this section) of a disease or disorder in family members of the individual; or

(D) Any request for, or receipt of, genetic services (as defined in paragraph (a)(4) of this section), or participation in clinical research which includes genetic services, by the individual or any family member of the individual.

(ii) The term genetic information does not include information about the sex or age of any individual.

(iii) The term genetic information includes—

(A) With respect to a pregnant woman (or a family member of the pregnant woman), genetic information of any fetus carried by the pregnant woman; and

(B) With respect to an individual (or a family member of the individual) who is utilizing an assisted reproductive technology, genetic information of any embryo legally held by the individual or family member.

(4) Genetic services means—

(i) A genetic test, as defined in paragraph (a)(5) of this section;

(ii) Genetic counseling (including obtaining, interpreting, or assessing genetic information); or

(iii) Genetic education.

(5)(i) Genetic test means an analysis of human DNA, RNA, chromosomes, proteins, or metabolites, if the analysis detects genotypes, mutations, or chromosomal changes. However, a genetic test does not include an analysis of proteins or metabolites that is directly related to a manifested disease, disorder, or pathological condition. Accordingly, a test to determine whether an individual has a BRCA1 or BRCA2 variant is a genetic test. Similarly, a test to determine whether an individual has a genetic variant associated with hereditary nonpolyposis colorectal cancer is a genetic test. However, an HIV test, complete blood count, cholesterol test, liver function test, or test for the presence of alcohol or drugs is not a genetic test.

(ii) The rules of this paragraph (a)(5) are illustrated by the following example:

Example. (i) Facts. Individual A is a newborn covered under a group health plan. A undergoes a phenylketonuria (PKU) screening, which measures the concentration of a metabolite, phenylalanine, in A’s blood. In PKU, a mutation occurs in the phenylalanine hydroxylase (PAH) gene which contains instructions for making the enzyme needed to break down the amino acid phenylalanine. Individuals with the mutation, who have a deficiency in the enzyme to
break down phenylalanine, have high concentrations of phenylalanine.

(ii) Conclusion. In this Example, the PKU screening is a genetic test with respect to A because the screening is an analysis of metabolites that detects a genetic mutation.

Example 2. (i) Facts. Individual B has a family member with colon cancer. One of them underwent genetic testing which detected a mutation in the MSH2 gene associated with hereditary nonpolyposis colorectal cancer (HNPCC). B’s physician, a health care professional with appropriate training and expertise in the field of medicine involved, has made a diagnosis that is not based principally on genetic information. Therefore, Huntington’s Disease by a health care professional with appropriate training and expertise for diagnosing Huntington’s Disease. The examination includes a clinical neurological exam. The results of the examination do not support a diagnosis of Huntington’s Disease.

(ii) Conclusion. In this Example 2, B has not been and could not reasonably be diagnosed with HNPCC. Thus, HNPCC is not manifested with respect to B.

Example 3. (i) Facts. Same facts as Example 2, except that B’s colonoscopy and subsequent test results indicate the presence of HNPCC. Based on the colonoscopy and subsequent test results, B’s physician makes a diagnosis of HNPCC.

(ii) Conclusion. In this Example 3, HNPCC is manifested with respect to B because a health care professional with appropriate training and expertise in the field of medicine involved has made a diagnosis that is not based principally on genetic information.

Example 4. (i) Facts. Individual C has a family member that has been diagnosed with Huntington’s Disease. A genetic test indicates that C has the Huntington’s Disease gene variant. At age 42, C begins suffering from occasional moodiness and disorientation, symptoms which are associated with Huntington’s Disease. C is examined by a neurologist (a physician with appropriate training and expertise for diagnosing Huntington’s Disease). The examination includes a clinical neurological exam. The results of the examination do not support a diagnosis of Huntington’s Disease.

(ii) Conclusion. In this Example 4, C is not and could not reasonably be diagnosed with Huntington’s Disease by a health care professional with appropriate training and expertise. Therefore, Huntington’s Disease is not manifested with respect to C.

Example 5. (i) Facts. Same facts as Example 4, except that C exhibits additional neurological and behavioral symptoms, and the results of the examination support a diagnosis of Huntington’s Disease with respect to C.

(ii) Conclusion. In this Example 5, C could reasonably be diagnosed with Huntington’s Disease by a health care professional with appropriate training and expertise. Therefore, Huntington’s Disease is manifested with respect to C.

(7) Underwriting purposes has the meaning given in paragraph (d)(1) of this section.

(b) No group-based discrimination based on genetic information—(1) In general. For purposes of this section, a group health plan must not adjust premium or contribution amounts for any employer, or any group of similarly situated individuals under the plan, on the basis of genetic information. For this purpose, “similarly situated individuals” are those described in §54.9802–1(d).

(2) Rule of construction. Nothing in paragraph (b)(1) of this section (or in paragraph (d)(1) or (d)(2) of this section) limits the ability of a group health plan to increase the premium for an employer or for a group of similarly situated individuals under the plan based on the manifestation of a
disease or disorder of an individual who is enrolled in the plan. In such a case, however, the manifestation of a disease or disorder in one individual cannot also be used as genetic information about other group members to further increase the premium for an employer or a group of similarly situated individuals under the plan.

(3) Examples. The rules of this paragraph (b) are illustrated by the following examples:

Example 1. (i) Facts. An employer sponsors a group health plan that provides coverage through a health insurance issuer. In order to determine the premium rate for the upcoming plan year, the issuer reviews the claims experience of individuals covered under the plan and other health status information of the individuals, including genetic information. The issuer finds that three individuals covered under the plan had unusually high claims experience. In addition, the issuer finds that the genetic information of two other individuals indicates the individuals have a higher probability of developing certain illnesses although the illnesses are not manifested at this time. The issuer quotes the plan a higher per-participant rate because of both the genetic information and the higher claims experience.

(ii) Conclusion. See Example 1 in 29 CFR 2590.702–1(b)(3) or 45 CFR 146.122(b)(3) for a conclusion that the issuer violates the provisions of 29 CFR 2590.702–1(b) or 45 CFR 146.122(b) similar to the requirements of this paragraph (b) because, by taking the likelihood that A’s children may develop poly-cystic kidney disease into account in computing the rate for the plan, the issuer adjusts the premium based on genetic information relating to a condition that has not been manifested in A’s children. However, the issuer does not violate the requirements of 29 CFR 2590.702–1(b) or 45 CFR 146.122(b) similar to the requirements of this paragraph (b) by increasing the premium based on A’s claims experience.

(c) Limitation on requesting or requiring genetic testing—(1) General rule. Except as otherwise provided in this paragraph (c), a group health plan must not request or require an individual or a family member of the individual to undergo a genetic test.

(2) Health care professional may recommend a genetic test. Nothing in paragraph (c)(1) of this section limits the authority of a health care professional who is providing health care services to an individual to request that the individual undergo a genetic test.

(3) Examples. The rules of paragraphs (c)(1) and (c)(2) of this section are illustrated by the following examples:

Example 1. (i) Facts. Individual A goes to a physician for a routine physical examination. The physician reviews A’s family medical history and A informs the physician that A’s mother has been diagnosed with Huntington’s Disease. The physician advises A that Huntington’s Disease is hereditary and recommends that A undergo a genetic test.

(ii) Conclusion. In this Example 1, the physician is a health care professional who is providing health care services to A. Therefore, the physician’s recommendation that A undergo the genetic test does not violate this paragraph (c).

Example 2. (i) Facts. Individual B is covered by a health maintenance organization (HMO). B is a child being treated for leukemia. B’s physician, who is employed by the HMO, is considering a treatment plan that includes six-mercaptopurine, a drug for treating leukemia in most children. However, the drug could be fatal if taken by a small percentage of children with a particular gene variant. B’s physician recommends that B undergo a genetic test to detect this variant before proceeding with this course of treatment.
(ii) Conclusion. In this Example 2, even though the physician is employed by the HMO, the physician is nonetheless a health care professional who is providing health care services to B. Therefore, the physician's recommendation that B undergo the genetic test does not violate this paragraph (c).

(4) Determination regarding payment—

(i) In general. As provided in this paragraph (c)(4), nothing in paragraph (c)(1) of this section precludes a plan from obtaining and using the results of a genetic test in making a determination regarding payment. For this purpose, “payment” has the meaning given such term in 45 CFR 164.501 of the privacy regulations issued under the Health Insurance Portability and Accountability Act. Thus, if a plan conditions payment for an item or service based on its medical appropriateness and the medical appropriateness of the item or service depends on the genetic makeup of a patient, then the plan is permitted to condition payment for the item or service on the outcome of a genetic test. The plan may also refuse payment if the patient does not undergo the genetic test.

(ii) Limitation. A plan is permitted to request only the minimum amount of information necessary to make a determination regarding payment. The minimum amount of information necessary is determined in accordance with the minimum necessary standard in 45 CFR 164.502(b) of the privacy regulations issued under the Health Insurance Portability and Accountability Act.

(iii) Examples. See paragraph (e) of this section for examples illustrating the rules of this paragraph (c)(4), as well as other provisions of this section.

(5) Research exception. Notwithstanding paragraph (c)(3) of this section, a plan may request, but not require, that a participant or beneficiary undergo a genetic test if all of the conditions of this paragraph (c)(5) are met:

(i) Research in accordance with Federal regulations and applicable State or local law or regulations. The plan makes the request pursuant to research, as defined in 45 CFR 46.102(d), that complies with 45 CFR Part 46 or equivalent Federal regulations, and any applicable State or local law or regulations for the protection of human subjects in research.

(ii) Written request for participation in research. The plan makes the request in writing, and the request clearly indicates to each participant or beneficiary (or, in the case of a minor child, to the legal guardian of the beneficiary) that—

(A) Compliance with the request is voluntary; and

(B) Noncompliance will have no effect on eligibility for benefits (as described in §54.9802–1(b)(1)) or premium or contribution amounts.

(iii) Prohibition on underwriting. No genetic information collected or acquired under this paragraph (c)(5) can be used for underwriting purposes (as described in paragraph (d)(1) of this section).

(iv) Notice to Federal agencies. The plan completes a copy of the “Notice of Research Exception under the Genetic Information Nondiscrimination Act” authorized by the Secretary and provides the notice to the address specified in the instructions thereto.

(d) Prohibitions on collection of genetic information—

(1) For underwriting purposes—

(i) General rule. A group health plan must not collect (as defined in paragraph (a)(1) of this section) genetic information for underwriting purposes. See paragraph (e) of this section for examples illustrating the rules of this paragraph (d)(1), as well as other provisions of this section.

(ii) Underwriting purposes defined. Subject to paragraph (d)(1)(iii) of this section, underwriting purposes means, with respect to any group health plan, or health insurance coverage offered in connection with a group health plan—

(A) Rules for, or determination of, eligibility (including enrollment and continued eligibility) for benefits under the plan or coverage as described in §54.9802–1(b)(1)(ii) (including changes in deductibles or other cost-sharing mechanisms in return for activities such as completing a health risk assessment or participating in a wellness program);

(B) The computation of premium or contribution amounts under the plan or coverage (including discounts, rebates, payments in kind, or other premium differential mechanisms in return for activities such as completing a
health risk assessment or participating in a wellness program):

(C) The application of any pre-existing condition exclusion under the plan or coverage; and

(D) Other activities related to the creation, renewal, or replacement of a contract of health insurance or health benefits.

(iii) Medical appropriateness. If an individual seeks a benefit under a group health plan, the plan may limit or exclude the benefit based on whether the benefit is medically appropriate, and the determination of whether the benefit is medically appropriate is not within the meaning of underwriting purposes. Accordingly, if an individual seeks a benefit under the plan and the plan conditions the benefit based on its medical appropriateness and the medical appropriateness of the benefit depends on genetic information of the individual, then the plan is permitted to condition the benefit on the genetic information. A plan is permitted to request only the minimum amount of genetic information necessary to determine medical appropriateness. The plan may deny the benefit if the patient does not provide the genetic information required to determine medical appropriateness. If an individual is not seeking a benefit, the medical appropriateness exception of this paragraph (d)(1)(iii) to the definition of underwriting purposes does not apply. See paragraph (e) of this section for examples illustrating the medical appropriateness provisions of this paragraph (d)(1)(iii), as well as other provisions of this section.

(2) Prior to or in connection with enrollment—(i) In general. A group health plan must not collect genetic information with respect to any individual prior to that individual’s effective date of coverage under that plan, nor in connection with the rules for eligibility (as defined in §54.9802-1(b)(1)(ii)) that apply to that individual. Whether or not an individual’s information is collected prior to that individual’s effective date of coverage is determined at the time of collection.

(ii) Incidental collection exception—(A) In general. If a group health plan obtains genetic information incidental to the collection of other information concerning any individual, the collection is not a violation of this paragraph (d)(2), as long as the collection is for underwriting purposes in violation of paragraph (d)(1) of this section.

(B) Limitation. The incidental collection exception of this paragraph (d)(2)(ii) does not apply in connection with any collection where it is reasonable to anticipate that health information will be received, unless the collection explicitly states that genetic information should not be provided.

(3) Examples. The rules of this paragraph (d) are illustrated by the following examples:

Example 1. (i) Facts. A group health plan provides a premium reduction to enrollees who complete a health risk assessment. The health risk assessment is requested to be completed after enrollment. Whether or not it is completed or what responses are given on it has no effect on an individual’s enrollment status, or on the enrollment status of members of the individual’s family. The health risk assessment includes questions about the individual’s family medical history.

(ii) Conclusion. In this Example 1, the health risk assessment includes a request for genetic information (that is, the individual’s family medical history). Because completing the health risk assessment results in a premium reduction, the request for genetic information is for underwriting purposes. Consequently, the request violates the prohibition on the collection of genetic information in paragraph (d)(1) of this section.

Example 2. (i) Facts. The same facts as Example 1, except there is no premium reduction or any other reward for completing the health risk assessment.

(ii) Conclusion. In this Example 2, the request is not for underwriting purposes, nor is it prior to or in connection with enrollment. Therefore, it does not violate the prohibition on the collection of genetic information in this paragraph (d).

Example 3. (i) Facts. A group health plan requests that enrollees complete a health risk assessment prior to enrollment, and includes questions about the individual’s family medical history. There is no reward or penalty for completing the health risk assessment.

(ii) Conclusion. In this Example 3, because the health risk assessment includes a request for genetic information (that is, the individual’s family medical history), and requests the information prior to enrollment, the request violates the prohibition on the collection of genetic information in paragraph (d)(2) of this section. Moreover, because it is a request for genetic information,
it is not an incidental collection under paragraph (d)(2)(ii) of this section.

Example 4. (i) Facts. The facts are the same as in Example 1, except there is no premium reduction or any other reward given for completion of the health risk assessment. However, certain people completing the health risk assessment may become eligible for additional benefits under the plan by being enrolled in a disease management program based on their answers to questions about family medical history. Other people may become eligible for the disease management program based solely on their answers to questions about their individual medical history.

(ii) Conclusion. In this Example 4, the request for information about an individual's family medical history could result in the individual being eligible for benefits for which the individual would not otherwise be eligible. Therefore, the questions about family medical history on the health risk assessment are a request for genetic information for underwriting purposes and are prohibited under this paragraph (d). Although the plan conditions eligibility for the disease management program based on determinations of medical appropriateness, the exception for determinations of medical appropriateness does not apply because the individual is not seeking benefits.

Example 5. (i) Facts. A group health plan requests enrollees to complete two distinct health risk assessments (HRAs) after and unrelated to enrollment. The first HRA instructs the individual to answer only for the individual and not for the individual's family. The first HRA does not ask about any genetic tests the individual has undergone or any genetic services the individual has received. The plan offers a reward for completing the first HRA. The second HRA asks about family medical history and the results of genetic tests the individual has undergone. The plan offers no reward for completing the second HRA and the instructions make clear that completion of the second HRA is wholly voluntary and will not affect the reward given for completion of the first HRA.

(ii) Conclusion. In this Example 5, no genetic information is collected in connection with the first HRA, which offers a reward, and no benefits or other rewards are conditioned on the request for genetic information in the second HRA. Consequently, the request for genetic information in the second HRA is not for underwriting purposes, and the two HRAs do not violate the prohibition on the collection of genetic information in this paragraph (d).

Example 6. (i) Facts. A group health plan waives its annual deductible for enrollees who complete an HRA. The HRA is requested to be completed after enrollment. Whether or not the HRA is completed or what responses are given on it has no effect on an individual's enrollment status, or on the enrollment status of members of the individual's family. The HRA does not include any direct questions about the individual's genetic information (including family medical history). However, the last question reads, "Is there anything else relevant to your health that you would like us to know or discuss with you?"

(ii) Conclusion. In this Example 6, the plan's request for medical information does not explicitly state that genetic information should not be provided. Therefore, any genetic information collected in response to the question is not within the incidental collection exception and is prohibited under this paragraph (d).

Example 7. (i) Facts. Same facts as Example 6, except that the last question goes on to state, "In answering this question, you should not include any genetic information. That is, please do not include any family medical history or any information related to genetic testing, genetic services, genetic counseling, or genetic diseases for which you believe you may be at risk."

(ii) Conclusion. In this Example 7, the plan's request for medical information explicitly states that genetic information should not be provided. Therefore, any genetic information collected in response to the question is within the incidental collection exception. However, the plan may not use any genetic information it obtains incidentally for underwriting purposes.

Example 8. (i) Facts. Issuer M acquires Issuer N's request for genetic information. However, the plan may not use any genetic information obtained incidentally for underwriting purposes.

(ii) Conclusion. In this Example 8, Issuer M's request for health information explicitly states that genetic information should not be provided. See Example 8 in 29 CFR 2590.702-1(d)(3) or 45 CFR 146.122(d)(3) for a conclusion that the collection of genetic information was within the incidental collection exception of 29 CFR 2590.702-1(d)(2)(ii) or 45 CFR 146.122(d)(ii) similar to the incidental exception of paragraph (d)(2)(ii) of this section. See Example 8 in 29 CFR 2590.702-1(d)(3) or 45 CFR 146.122(d)(3) also for a caveat that M may not use the genetic information it obtained incidentally for underwriting purposes.

(e) Examples regarding determinations of medical appropriateness. The application of the rules of paragraphs (c) and
Example 1. (i) Facts. Individual A’s group health plan covers genetic testing for celiac disease for individuals who have family members with this condition. After A’s son is diagnosed with celiac disease, A undergoes a genetic test and promptly submits a claim for the test to A’s issuer for reimbursement. The issuer asks A to provide the results of the genetic test before the claim is paid.

(ii) Conclusion. See Example 1 in 29 CFR 2590.702-1(e) or 45 CFR 146.122(e) for a conclusion under the rules of paragraph (c)(4) of 29 CFR 2590.702-1 or 45 CFR 146.122 similar to the rules of paragraph (c)(4) of this section that the issuer is permitted to request only the minimum amount of information necessary to make a decision regarding payment. Because the results of the test are not necessary for the issuer to make a decision regarding the payment of A’s claim, the conclusion in Example 1 in 29 CFR 2590.702-1(e) or 45 CFR 146.122(e) concludes that the issuer’s request for the results of the genetic test violates paragraph (c) of 29 CFR 2590.702-1 or 45 CFR 146.122 similar to paragraph (c) of this section.

Example 2. (i) Facts. Individual B’s group health plan covers a yearly mammogram for participants and beneficiaries starting at age 40, or at age 30 for those with increased risk for breast cancer, including individuals with BRCA1 or BRCA2 gene mutations. B is 33 years old and has the BRCA2 mutation. B undergoes a mammogram and promptly submits a claim to B’s plan for reimbursement. Following an established policy, the plan sends out a notice to all participants that describes the diabetes disease management program and explains the terms for eligibility. Individuals interested in enrolling in the program are advised to contact the plan to demonstrate that they have diabetes or that they are at risk for diabetes. For individuals who do not currently have diabetes, genetic information may be used to demonstrate that an individual is at risk.

(ii) Conclusion. In this Example 2, the plan does not violate paragraph (c) or (d) of this section. Under paragraph (c), the plan is permitted to request and use the results of a genetic test to make a determination regarding medical appropriateness for the individual and whether the disease management program is medically appropriate for the individual and only requests the minimum amount of information necessary. The genetic information is not used for underwriting purposes.

Example 3. (i) Facts. Individual C was previously diagnosed with and treated for breast cancer, which is currently in remission. In accordance with the recommendation of C’s physician, C has been taking a regular dose of tamoxifen to help prevent a recurrence. C’s group health plan adopts a new policy requiring patients taking tamoxifen to undergo a genetic test to ensure that tamoxifen is medically appropriate for their genetic makeup. In accordance with, at the time, the latest scientific research, tamoxifen is not helpful in up to 7 percent of breast cancer patients, those with certain variations of the gene for making the CYP2D6 enzyme. If a patient has a gene variant making tamoxifen not medically appropriate, the plan does not pay for the tamoxifen prescription.

(ii) Conclusion. In this Example 3, the plan does not violate paragraph (c) of this section if it conditions future payments for the tamoxifen prescription on C’s undergoing a genetic test to determine what genetic markers C has for making the CYP2D6 enzyme. Nor does the plan violate paragraph (c) of this section if the plan refuses future payment if the results of the genetic test indicate that tamoxifen is not medically appropriate for C.

Example 4. (i) Facts. A group health plan offers a diabetes disease management program to all similarly situated individuals for whom it is medically appropriate based on whether the individuals have or are at risk for diabetes. The program offers enhanced benefits related only to diabetes for individuals who qualify for the program. The plan sends out a notice to all participants that describes the diabetes disease management program and explains the terms for eligibility. Individuals interested in enrolling in the program are advised to contact the plan to demonstrate that they have diabetes or that they are at risk for diabetes. For individuals who do not currently have diabetes, genetic information may be used to demonstrate that an individual is at risk.

(ii) Conclusion. In this Example 4, the plan may condition benefits under the disease management program upon a showing by an individual that the individual is at risk for diabetes, even if such showing may involve genetic information, provided that the plan requests genetic information only when necessary to make a determination regarding whether the disease management program is medically appropriate for the individual and only requests the minimum amount of information necessary to make that determination.

Example 5. (i) Facts. Same facts as Example 4, except that the plan includes a questionnaire that asks about the occurrence of diabetes in members of the individual’s family...
§54.9811–1 Standards relating to benefits for mothers and newborns.

(a) Hospital length of stay—(1) General rule. Except as provided in paragraph (a)(5) of this section, a group health plan that provides benefits for a hospital length of stay in connection with childbirth for a mother or her newborn may not restrict benefits for the stay to less than—

(i) 48 hours following a vaginal delivery; or

(ii) 96 hours following a delivery by cesarean section.

(2) When stay begins—(i) Delivery in a hospital. If delivery occurs in a hospital, the hospital length of stay for the mother or newborn child begins at the time of delivery (or in the case of multiple births, at the time of the last delivery).

(ii) Delivery outside a hospital. If delivery occurs outside a hospital, the hospital length of stay begins at the time the mother or newborn is admitted as a hospital inpatient in connection with childbirth. The determination of whether an admission is in connection with childbirth is a medical decision to be made by the attending provider.

(3) Examples. The rules of paragraphs (a)(1) and (2) of this section are illustrated by the following examples. In each example, the group health plan provides benefits for hospital lengths of stay in connection with childbirth and is subject to the requirements of this section, as follows:

Example 1. (i) Facts. A pregnant woman covered under a group health plan goes into labor and is admitted to the hospital at 10 p.m. on June 11. She gives birth by vaginal delivery at 6 a.m. on June 12.

(ii) Conclusion. In this Example 1, the 48-hour period described in paragraph (a)(1)(i) of this section ends at 6 a.m. on June 14.

Example 2. (i) Facts. A woman covered under a group health plan gives birth at home by vaginal delivery. After the delivery, the woman begins bleeding excessively in connection with the childbirth and is admitted to the hospital for treatment of the excessive bleeding at 7 p.m. on October 1.

(ii) Conclusion. In this Example 2, the 48-hour period described in paragraph (a)(1)(i) of this section ends at 7 p.m. on October 3.

Example 3. (i) Facts. A woman covered under a group health plan gives birth by vaginal delivery at home. The child later develops pneumonia and is admitted to the hospital. The attending provider determines that the admission is not in connection with childbirth.

(ii) Conclusion. In this Example 3, the hospital length-of-stay requirements of this section do not apply to the child’s admission to the hospital because the admission is not in connection with childbirth.

(4) Authorization not required—(i) In general. A plan may not require that a physician or other health care provider obtain authorization from the plan, or from a health insurance issuer offering health insurance coverage under the plan, for prescribing the hospital length of stay specified in paragraph (a)(1) of this section. (See also paragraphs (b)(2) and (c)(3) of this section for rules and examples regarding other authorization and certain notice requirements.)