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Chapter 5 Conclusions and Recommendations

An ethical research process pursues its scientific aims without compromising the rights and welfare of human subjects. Achieving such a balance is a particular challenge in rapidly advancing fields, of which human genetics (and molecular biology generally) is currently a prime example. In such situations, the tantalizing potential for generating major advances can make research activities seem especially important and compelling. At the same time, the novelty of the field can mean that potential harms to subjects are poorly understood or seem largely speculative. This is particularly true of non-physical injuries to subjects' interests such as those that can arise in research on previously collected human biological materials when investigators do not physically interact with the persons whose tissues, cells, or DNA they are studying.

Research sponsors, investigators, and IRBs thus need to exercise great care and sensitivity in applying professional guidelines and government regulations to protect subjects whose biological specimens are used in research. Properly interpreted and modestly modified, present federal regulations can protect subjects' rights and interests while at the same time permitting well designed research to go forward using specimens already in storage as well as those newly collected by investigators. The interests of subjects and those of researchers are not fundamentally in conflict, for appropriate protection of subjects provides the reassurance needed if individuals are to continue to make their tissue, blood, or DNA available for research. Indeed, public confidence in the ethics and integrity of the research process translates into popular support for research generally.

For most people, the central issue in research using human tissues and cells is the harm that can occur when private information about present or future health status—often previously unknown even to the subject—is revealed. One simple protection for subjects would be to render anonymous all human biological samples used in research. That solution would, however,

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1 seriously curtail many investigations. Instead, the protection of human subjects should take
2 account of the great value for many studies using human biological materials of having access to
3 a certain amount of personal and clinical data about the persons from whom specimens were
4 obtained. That is, the policies and guidelines governing human subjects research ought—under
5 certain circumstances—to permit investigators to have access to sufficient identifying information
6 to enable them to gather necessary data about subjects. Assuming that adequate protections
7 (including informed, voluntary consent) are present, such information-gathering could include
8 on-going collection of medical record data and even requests for subjects to undergo tests to
9 provide additional research data. In some cases, it will even be acceptable for investigators to
10 convey information about research results back to the persons whose samples have been studied.
11 Where identifying information exists, however, a well-developed system of protections must be
12 implemented to ensure that risks are minimized and the interests of sample sources are protected.

13

14 Finally, any system of regulation is most likely to achieve its goals if it is as clear and
15 simple as possible. This is especially true here because the federal protections for research
16 subjects depend on investigators identifying the involvement of human subjects in their studies
17 and initiating the institutional review of their protocols. Thus, one reason to modify regulations is
18 to make clearer their description of which protocols are subject to what sorts of prior review;
19 likewise, illustrations and explanations may be useful to clarify how the regulations apply to
20 novel or complicated fields, such as genetic studies using human biological materials.

21

22 How well does the existing Federal Policy for the Protection of Human Subjects (the so-
23 called “Common Rule,” codified at 45 CFR Part 46) meet these objectives? Specifically, does it
24 provide clear direction to research sponsors, investigators, IRBs, and others regarding how to
25 conduct research using human biological materials in an ethically sound manner? Not entirely. In
26 some cases present regulatory language provides ambiguous guidance for research using human
27 biological material, though the language can be retained if correctly interpreted. For example,

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1 confusion about the intended meaning of terms such as “existing samples,” “publicly available,”
2 and “minimal risk” has stymied investigators and IRB members.¹ Beyond these ambiguities,
3 certain parts of current regulations do not adequately ensure the ethical use of human biological
4 materials in research, but require some modification. This Chapter provides interpretations of
5 several important concepts and terms in the Common Rule and recommends ways both to
6 strengthen and clarify the regulations and to make their implementation more consistent.

7
8 The goals of these recommendations are to: (1) address perceived difficulties in the
9 interpretation of federal regulations, and in the language of some professional organizations; 2)
10 ensure that research involving human biological materials will continue to benefit from
11 appropriate oversight and IRB review, the additional burdens of which are kept to a minimum; 3)
12 provide investigators and IRBs with clear guidance regarding the use of human biological
13 materials in research, particularly with regard to informed consent; 4) provide a coherent public
14 policy process framework for research in this area that will endure for many years and be
15 responsive to new developments in science; and 5) provide the public (including potential
16 research subjects) with increased confidence in the research activity. To accomplish these goals,
17 NBAC makes 26 recommendations in the following areas:

- 18 • adequacy and interpretation of existing federal policies for the protection of human subjects
- 19 • informed consent
- 20 • waiver of consent
- 21 • reporting of research results to subjects
- 22 • consideration of potential harms to others
- 23 • publication and dissemination of study results
- 24 • professional education and responsibilities
- 25 • federal and state legislation on medical record privacy

¹ See testimony of Mary-Claire King, Ph.D., before NBAC, July 1998, Portland, Oregon.

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Research Activities Beyond Clinical Care

In order to trigger the regulations, an activity must be considered “research,” as opposed to a clinical intervention. The current regulations do not apply to purely clinical uses of human biological materials, or to other activities such as quality control procedures or teaching. Rather, they apply to *research* defined as “a systematic investigation designed to develop or contribute to generalizable knowledge” (45 CFR 46.102(d)). *Research on stored materials undertaken solely as part of a clinical intervention—as when a pathologist assesses a biopsied specimen to confirm a diagnosis—falls outside the purview of this report, although activities that have both a clinical and a research component are covered.* Thus, any study conducted on samples left over from a clinical intervention is subject to the federal research regulations, if the investigator or the investigator’s institution is subject to those regulations or if the institution has voluntarily agreed not to supply samples for research without following the federal regulations.² As investigators make greater use of human biological materials in a wide range of research, it thus becomes important that the expectations of the federal regulations be communicated to—and acted upon by—specimen repositories in a timely fashion in collecting specimens for clinical purposes.

Adequacy and Interpretation of the Existing Federal Policy for the Protection of Human Subjects

In the context of studies using human biological materials, the lack of clarity regarding several regulatory terms means that they do not provide the guidance needed by investigators, IRBs, and others. These terms include: “existing and publicly available,” “identifiable,” “minimal

²The protections provided by the Common Rule currently apply only to: 1) research conducted or funded by one of the 17 agencies that have agreed to be subject to the Common Rule or by any other federal agency that has promulgated its own set of human subjects research rules; or 2) research conducted at an institution that has provided in its “assurance” with the federal government that all research with human subjects conducted as the institution will be governed by the federal regulations whether or not the research is federally sponsored. In addition, the FDA 2) regulates human subjects research involving an investigational new drug, device, or biologic.

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1 risk,” “rights and welfare,” and “practicable” .

2
3 **“Existing and Publicly Available”**

4 Under two conditions, research with human biological materials may be exempt from the
5 Common Rule, namely when:

- 6 1) the samples are existing and publicly available; or
7 2) the samples are existing and information is recorded by the investigator in such a
8 manner that subjects cannot be identified, directly or through identifiers linked to the
9 subjects (45 CFR 46.101 (b) (4)).

10
11 Furthermore, since current federal regulations define a human subject as a “living
12 individual,” research using stored specimens from people who have died would not come with
13 the regulatory protection for human subjects.³

14
15 Regarding the first exemption (that the samples are existing and publicly available), the
16 Office for Protection from Research Risk (OPRR) interprets the term “existing” to mean any
17 materials that are already collected—that is, “on the shelf”—at the time the research is initiated,
18 whether collected for previous research or nonresearch purposes.⁴ “Existing” thus contrasts with
19 samples that are to be collected at a later date as a part of the research protocol in question.

20
21 The second criterion in the first exemption—the requirement that samples be “publicly
22 available”—is more problematic. In response to a request for clarification, OPRR defined
23 publicly available to mean that “unrestricted access on demand (*i.e.*, unrestricted availability

³ See 45 CFR 46.102 (f). If the source of the sample is deceased, then according to the regulations, there is no human subject and the regulations do not apply. As discussed below, circumstances may exist in which research on samples of deceased individuals has implications for living relatives; if the findings were attached in some way to these relatives, they might be considered human subjects, which could trigger the federal regulations.

⁴ See, e.g., IRB Guidebook, pp. xxx.

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1 subject only to limited quantities and/or related costs) may be considered a reasonable basis for
2 claiming ‘publicly available’.’⁵ Yet this interpretation provides minimal guidance as it leaves
3 unclear which “public” is the relevant (*e.g.*, the general public, the scientific community) and
4 whether “available” is the same as “accessible”.

5
6 Large repositories, often cited as examples of “public collections” have in place “strict
7 policies to ensure that cultures are distributed only to qualified organizations and researchers with
8 legitimate and justifiable scientific uses for these materials.”⁶ Thus, the biological materials are
9 available not to *anyone*, but are, in general, restricted to those who have a legitimate research
10 interest in their use and presumably possess the capabilities to perform sophisticated scientific
11 studies that can reveal biological information about that sample or even health-related
12 information about the person from whom it came. Moreover, some newer DNA databases (*e.g.*,
13 those associated with the federally funded Human Genome Project) are organized on the
14 assumption that such information *should be* available to any scientist wanting to investigate the
15 basic structure or function of DNA. For example, the National Human Genome Research
16 Institute implements a policy requiring that primary human genomic sequence data be rapidly
17 released (within 24 hours of generation).

18
19 Thus, although collections might be widely available to the research community, and
20 appropriately so, it appears that they are infrequently available to any member of the public.
21 While the interests of the people who have provided these specimens are best protected by
22 restricting access to these materials to qualified researchers, the fact that such researchers can
23 readily access the specimens does not make them “publicly available” as that term is commonly
24 understood. The reason for exempting publicly available data from the purview of the Common
25 Rule is that people have no expectation of privacy about information that anyone can find and

⁵ Personal communication from OPRR Director, Dr. Gary Ellis, August 25, 1998.

⁶ American Type Culture Collection (ATCC), <http://www.atcc.org/>

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1 that any harm that may be associated with the disclosure of such information has already
2 occurred and should, in any case, be taken up with those who collected the data and made them
3 public.

4
5 **Recommendation 1:**

6
7 **Current regulations state that research using *existing* human biological materials**
8 **that are publicly available is exempt from IRB review (45 CFR 46.101(b)(4)). Few, if**
9 **any, collections of human biological materials are publicly available. Therefore, the**
10 **Office for Protection from Research Risks should make clear in its guidance that in**
11 **most cases this exemption does not apply to research using human biological**
12 **materials.**

13
14 It should be noted that the current regulatory policy makes sense in the context for which
15 it was first created—such as a social or behavioral scientist making use of information about
16 people that can be found in directories or newspapers or observed in recordings made of their
17 conduct in public settings. But the exemption would contradict the whole purpose of human
18 subjects protection were it applied to new biological analyses of stored human tissues or cells
19 because the information that may emerge from such a process is not in any genuine sense
20 “existing” much less “publicly available.” Furthermore, since the first exemption would
21 encompass data that are identified with individual human subjects, applying that exemption to
22 human specimens in repositories to which researchers have easy access would render moot the
23 second exemption for studies using existing samples, which limits investigators to information
24 that cannot be linked to the subjects.

25
26 Thus, while the accessibility of specimens is an important consideration in specifying
27 appropriate levels of protection, more important considerations include: 1) whether the

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1 specimens are stored with codes, links, or identifiers; 2) whether identifiable samples (coded or
2 identified) are delivered to investigators for study; and 3) whether the repositories or retainers of
3 the specimens require any assurance that the research will be conducted in a manner that will
4 protect the rights and interests of the sources.⁷

The “Identifiability” of Samples and the Applicability of the Common Rule

7 The second criterion for exemption from the Common Rule (that the samples are existing
8 and information is recorded by the investigator in such a manner that subjects cannot be
9 identified, directly or through identifiers linked to the subjects) reflects an underlying premise of
10 the federal regulations, namely that protection is needed when research is conducted with *human*
11 *subjects*. The regulations define a human subject as “a living individual about whom an
12 investigator conducting research obtains: (a) data through intervention or interaction with the
13 individual, or (b) identifiable private information.”⁸ Section 46.102(f)(2) defines “identifiable” to
14 mean “the identity of the subject is or may readily be ascertained by the investigator
15 or...associated with the information.” OPRR interprets “identifiable” to include specimens with
16 codes that, with the cooperation of others, could be broken in order to reveal the name of the
17 tissue source.⁹

19 The academic and professional literature on the research use of human biological
20 materials has utilized a variety of terms to describe the identifiability of research samples. Part of
21 the confusion about the interpretation of the term “identifiable” arises from the fact that people
22 sometimes refer to the state of the information attached to the biological material in the repository

⁷ In reviewing the policies and procedures of several repositories, the Commission found that some require that investigators provide a statement of their research intent and an assurance of compliance with the Common Rule, but it is not clear that this practice is widespread, especially among smaller, more informal tissue collections.

⁸ 45 CFR 46.102(f)(1)&(2).

⁹ IRB GUIDEBOOK, pp. 2-9.

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1 (i.e., the **specimen**) and sometimes refer to the material (i.e., the **sample**) and the accompanying
2 information that is provided to the researcher. For example, the specimen might be identified in
3 the repository but no identifying information is forwarded with the research sample sent to the
4 researcher. This distinction is of considerable importance because the potential for both benefit
5 and harm is greater when the sample is directly or easily linked to the person who provided the
6 specimen, placing the burden of protection in different places, depending on who has access to
7 the information (e.g., the researcher or the pathologist, or both). If samples are identifiable, the
8 potential exists for the investigator or a third party (e.g., insurer, employer) to contact the subject
9 or act in some way that might affect the subject. For example, an investigator might want to
10 contact an individual to gather more medical information, obtain consent for additional or
11 different uses of the sample, inform them about the results of the study, or communicate findings
12 that might be of clinical significance to that individual.

13
14 NBAC adopted the following definitions regarding the diverse status of human biological
15 materials, depending on whether they are sitting in storage in a repository, or whether some of
16 the material from a repository has been selected for research purposes.

17
18 **Repository collections** of human biological materials (i.e., specimens) are one of two
19 types:

- 20 1. **Unidentified specimens** are those for which identifiable personal information was
21 not collected or, if once collected, is not maintained and cannot be retrieved by the
22 repository.
- 23 2. **Identified specimens** are those linked to personal information, such that the person
24 from whom the material was obtained could be identified by name, patient numbers,
25 or clear pedigree location (i.e., their relationship to a family member, whose identity is
26 known).

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1 **Research samples** are the collections of human biological materials provided to
2 investigators by repositories. Such materials are of at least four types, which are differentiated by
3 the amount of information that is conveyed to the investigator about the person from whom the
4 sample comes. NBAC defines the different types as follows:

- 5
6 1. **Unidentified samples**—sometimes termed “anonymous”—are those supplied by
7 repositories from an unidentified collection of human biological specimens.
- 8 2. **Unlinked samples**—sometimes termed “anonymized”—are those supplied by
9 repositories from identified human biological specimens without identifiers or codes
10 such that the ability to identify particular individuals via clinical or demographic
11 information supplied with the sample, or biological information derived from the
12 research that would be extremely difficult for the investigator, the repository, or a
13 third party.
- 14 3. **Coded samples**—sometimes termed “linked” or “identifiable”—are those supplied
15 by repositories from identified specimens with a code rather than a name or any other
16 personal identifier such as a patient number, where the repository (or its agent) retains
17 information linking the code to particular human specimens or where the extent of the
18 clinical or demographic information provided with the sample is sufficient that the
19 investigator, the repository, or a third party could link the biological information
20 derived from the research with material from a particular person or a very small group
21 of identifiable persons.
- 22 4. **Identified samples** are those supplied by repositories from identified specimens with
23 a personal identifier (such as a name or patient number) sufficient to allow the
24 biological information derived from the research to be linked directly, by the
25 researcher, with the particular person from whom the material was obtained.

26
27 For the purposes of interpreting and applying the regulations, NBAC aggregates these

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1 four groups into two categories: 1) *unidentifiable samples*, which are either unidentified or
2 unlinked (categories 1 and 2 above); and 2) *identifiable samples*, either coded or identified
3 (categories 3 and 4 above). The recommended protections required within each category are the
4 same.

6 ***Unidentifiable Samples***

7 As mentioned above, within the “unidentifiable” category are two subcategories: 1)
8 unidentified samples; and 2) unlinked samples. Unidentified samples have no data (even as
9 specimens in the repository) linking them to an individual and, therefore, no one has the ability to
10 determine the identity of the source of the specimen. Such samples are completely anonymous.
11 In other cases, the samples may be “unlinked” or “anonymized,” that is, the specimens from
12 which the samples are derived retain identifiers but the samples are forwarded to a researcher
13 without any identifiers or codes.

15 **Recommendation 2:**

17 **Research conducted on unidentified samples, whether taken from specimens stored**
18 **without personal identifiers or those rendered unidentifiable by someone**
19 **independent of the investigator, does not involve 'human subjects' and hence is not**
20 **subject to the requirements of the Common Rule. The Office for Protection from**
21 **Research Risks should issue appropriate guidance for investigators and**
22 **Institutional Review Boards on these definitions of identifiability, or, if deemed**
23 **necessary, modify the language of the regulations (“the Common Rule”).**

24
25 Several repositories keep a record of the persons from whom the samples came so that
26 the repository can track that a sample was sent to a clinician or researcher. Such samples may be
27 numbered in such a way so that even the repository cannot link the sample to its source. Or,

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1 samples might be numbered in such a way that the repository can track that a sample was sent
2 forward but if the investigator were to come back to the repository and ask for additional material
3 or clinical information specific to that source the repository could not match the request with a
4 specific specimen. At best, the repository could send the investigator a duplicate set of the initial
5 “batch” of samples, but again with no linking data. There might be some rare cases in which the
6 sample size is so small and the findings so unique that it would be relatively easy to identify
7 individuals even if their samples were unlinked. Investigators and repositories should give these
8 situations careful scrutiny to reduce the chance that persons could be identified. In such
9 instances, it may be more appropriate to use only unidentified (not merely “unlinked”) samples,
10 increase the sample size, or even consider the samples to be identifiable rather than
11 unidentifiable.

12

13 When researchers use unidentified and unlinked samples, contact of the source by the
14 researcher is extremely difficult. According to the federal regulations, research using existing
15 samples of this type is exempt from IRB review. The justification for this regulation appears to
16 be that since it is not possible to contact the sources to ask their permission for any specific uses
17 or to gain consent, and because the potential for harm effectively disappears due to lack of
18 identifiability, no special restrictions of the use of such unidentifiable samples should apply.

19

20 Although this seems quite reasonable at first, some controversy remains in the case of
21 samples that have been rendered unidentifiable before being sent on to the investigator. Some
22 might consider it ethically problematic that by having identifiers stripped, the investigator loses
23 the opportunity to obtain consent, since further recontact would be prevented. In addition, it is
24 incorrect to assume that because the sources cannot be identified they cannot be harmed or
25 wronged. There are some interests of the sample sources that may be harmed even if the sources
26 are not completely identifiable, and there may be some interests of others at risk as well. For
27 example, there might be group or family interests that could be revealed or placed at risk because

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1 of research done on a class of similar, albeit individually unidentifiable, samples. Individuals have
2 an interest in avoiding uses of their tissue that they regard as impermissible or objectionable on
3 moral grounds. Thus, were their samples to be used in research that they would find
4 objectionable then it is possible that some individuals could be wronged, if not harmed. NBAC
5 recognizes these concerns as valid but not sufficiently substantial to restrict further use of such
6 samples.

7
8 Because the samples are not linkable to individuals, some of the most important interests
9 that weigh in favor of restricted access do not apply. If the individual cannot be identified, then
10 there is little or no risk of insurance or employment discrimination, stigma, adverse psychological
11 reactions, or familial conflict. So to that extent, the case for not allowing use of unidentifiable
12 stored samples is significantly weakened. The possibility remains that research findings might
13 still result in potential harms to groups or classes of individuals (e.g., loss of health insurance
14 coverage for individuals found to share a particular trait or characteristic). Although the current
15 regulations do not require investigators to consider such risks to groups, good practice might, in
16 some cases, warrant an effort to minimize risks to others through consultation with relevant
17 groups, alterations in research design, or greater care in the manner in which research results are
18 reported. (See also Recommendations 19 and 20.)

19
20 Given the importance of society's interest in treating disease and developing new
21 therapies, a policy that severely restricted research access to these unidentifiable samples would
22 severely hamper research and could waste a valuable research resource.

23 24 ***Identifiable Samples***

25 Within the "identifiable" category are two subcategories: 1) coded samples; and 2)
26 identified samples (i.e., where the sample source is expressly identified to the investigator).
27 Within the first category there may be a distinction between the information provided to the

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1 investigator and that held by the repository. For example, the samples might be encoded in such
2 a way that the investigator cannot identify the sample source but the entity storing the sample,
3 such as a pathologist or DNA bank, can link the sample source to the specimen sent to the
4 investigator. Thus, the code could be broken if desired. Although identifying the source may be
5 more difficult in this latter scenario, NBAC considers these samples to be identifiable, because
6 the possibility of linkage remains, elevating the potential for harm. The ease of identifying the
7 source is part of the calculus in determining the overall level of risk posed by the research. This
8 matter is discussed later.

9
10 Previous guidelines and reports (see Chapter 3) have categorized samples by the
11 conditions under which they are stored (with or without identifiers). Current federal regulations
12 permit researchers to take existing samples, render them anonymous by removing identifiers, and
13 then use them in research without seeking consent. It is apparent to NBAC that some
14 investigators incorrectly interpret the regulations to mean that as long as **they** do not know the
15 identity of the sample source, even if the sample is coded (linked), the research is exempt from
16 IRB review. The issue of identifiability is further confounded by the researcher's growing ability
17 to identify the source (even when unidentified) because of the possibility that DNA analysis will
18 permit matching of samples with individuals. NBAC concluded that the policy would better
19 protect human subjects, while still preserving the scientific value of the samples, if someone
20 independent of the investigator coded the samples or rendered them unidentifiable, for example
21 the repository, an encryption service, or someone at the research institution who is not directly
22 involved in the conduct of the research in question.

23
24 **Recommendation 3:**

25
26 **Research conducted on human biological materials that are linked to information**
27 **that could identify the individuals from whom they were obtained, even through a**

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1 **code, is subject to the process of review and approval specified by the Common Rule**
2 **(45 CFR 46.101(b)(4).**

3
4 NBAC recognizes that there may be costs associated with this requirement. Thus, any
5 costs incurred by the investigator to satisfy this requirement should be considered by the funding
6 agency a valid and reimbursable expense.

7
8 NBAC does not believe that these interpretations of the criteria for exemption and review
9 will impede research. In fact, some repositories already have in place these protections and many
10 investigators voluntarily elect to have repositories strip identifiers before samples are sent forward
11 to their laboratories. These interpretations will ensure that research conducted on identifiable
12 samples, even if widely or publicly available, will be subject to the federal policy of protections.
13 Repositories and IRBs share responsibility with investigators to ensure that research is designed
14 and conducted in a manner that appropriately protects human subjects from unnecessary harms.

15
16 **Recommendation 4:**

17
18 **Before releasing identifiable samples from its collection, a repository should require**
19 **that the investigator requesting the samples either: a) provide documentation from**
20 **the investigator’s Institutional Review Board that the research will be conducted in**
21 **compliance with applicable federal regulations; or b) explain in writing why the**
22 **research is not subject to those regulations.**

23
24 **Recommendation 5:**

25
26 **When reviewing and approving a protocol for research on human biological**
27 **materials, Institutional Review Boards should require the investigator to set forth:**

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- 1 **a) a thorough justification of the research design, including a description of**
- 2 **procedures used to minimize risk to subjects;**
- 3 **b) a full description of the process by which samples will be obtained from**
- 4 **repositories;**
- 5 **c) any plans to access the medical records of the subjects; and**
- 6 **d) a full description of the mechanisms that will be used to maximize the protection**
- 7 **against inadvertent release of confidential information.**

9 **Obtaining Informed Consent**

10
11 Specimens that already exist in storage at the time the research is proposed may have been
12 collected under a variety of conditions (e.g., in a clinical setting or as part of an experimental
13 protocol). In some instances, individuals make informed choices about how their sample should
14 be used subsequent to its original research or clinical use. In other cases, for a variety of reasons,
15 individuals may not fully understand or may have not been given the opportunity to carefully
16 consider and decide how their sample may be used in the future. When research is contemplated
17 using existing samples, the expressed wishes of the individuals who provided the material must
18 be respected. Where consent documents exist, they may indicate whether individuals wanted
19 their sample to be used in future research, and in some instances the specific type of research.

20
21 IRBs should use the following criteria to evaluate the applicability of such documents to
22 the proposed research:

- 23 • Does the language or context of the consent form indicate that the source was interested in
24 aiding the type of research being proposed?
- 25 • If the person consented to the sample being used in unspecified future studies, is that consent
26 adequate for the type of research being planned, given the circumstances under which the
27 sample was collected (e.g., whether the sample was requested by a treating physician,

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1 whether the consent form offered alternatives to allowing the sample to be used in future
2 studies)?

3
4 In some cases an IRB may determine that an existing consent form permitting unspecified
5 future uses to be sufficient. For example, Clayton and colleagues argue that, “[e]ven in the
6 absence of specific language about DNA testing, it may be appropriate to infer consent if the
7 source wished for the sample to be used to determine why his or her family had a particular
8 inherited disorder (1995).” In such cases, investigators should consider informing subjects that
9 research is occurring and in certain cases also give them the opportunity to “opt out.” Rarely,
10 however, does the language in typical operative and hospital admission consent forms provide an
11 adequate basis for inferring consent to future research.

12
13 This policy provides that provides significant protection for sources, recognizes that their
14 samples may have been collected without adequate disclosure, yet provides them the opportunity
15 to participate in research. When the IRB determines existing consent documents to be inadequate
16 and where the existing sample is identifiable, the individual should be contacted, offered the
17 option of consenting to the specific proposed protocol, and further offered the option of deciding
18 how the sample may be used in the future.

19
20 As in the case with research in which new samples are obtained, individuals should be
21 provided with relevant information to assist them in making a decision about participation in
22 research. Federal human subjects regulations list the basic elements of informed consent which,
23 of course, apply also when consent is requested for the use of existing samples (45 CFR
24 46.116[a]). The following points are especially relevant here:

25
26 a) The risks and benefits of participation in the proposed study along with a discussion of
27 the possible consequences of consenting to future identifiable uses of their sample.

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- 1 b) The extent, if any, to which confidentiality will be maintained. (Investigators are
- 2 encouraged to seek certificates of confidentiality, when appropriate.)
- 3 c) Under what circumstances, if any, subjects will be re-contacted.
- 4 d) An indication that if subjects choose to have their sample rendered unidentifiable they
- 5 cannot be given specific information about findings related to their samples.

6

7 The rationale for including the option of authorization for future research use of existing

8 samples rather than mere disclosure that the sample may be used for a wide range of purposes is

9 that in most cases existing samples will have been collected without disclosure. Allowing

10 persons (whose previously collected samples are identifiable) to choose either to authorize future

11 research use or to have their samples rendered unidentifiable for future uses can be viewed as an

12 effort to repair this deficiency. Even if such authorization bears only a remote resemblance to

13 genuine informed consent, it can serve as an expression of respect for persons in the context of

14 proposed uses for existing samples. Simply to disclose to persons now that the sample already

15 taken from them may be used for purposes of which they had no idea at the time of collection is

16 not adequate.¹⁰

17

18 **Recommendation 6:**

19

20 **When conducting research on samples obtained prior to the release of this report,**

21 **general releases for research given in conjunction with a clinical or surgical procedure**

22 **must not be presumed to cover all types of research over an indefinite period of time.**

23 **Investigators and Institutional Review Boards should review existing consent**

24 **documents to determine whether the subjects anticipated and agreed to participate in**

25 **the type of research proposed. If the existing documents are inadequate, and consent**

¹⁰ Elsewhere, NBAC has discussed the issue of prospective authorization and found that under some circumstances it is an important method of respecting individual choices (see NBAC, *Research Involving Persons with Mental Disorders That May Affect Decisionmaking Capacity*, 1998, p.61). NBAC does not regard prospective authorization

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1 **cannot be waived, the investigator must obtain informed consent from the subjects for**
2 **the current research. (See Recommendations xx for waiver of consent.)**

3
4 Whether obtaining consent to the research use of human biological materials in a research
5 or clinical setting, and whether the consent is new or renewed, efforts should be made to be as
6 explicit as possible about the uses to which the material might be put and whether there is a
7 possibility that such research might be done in such a way that the individual could be identified.
8 Obviously, different conditions will exist for different protocols, in different settings, and among
9 individuals. NBAC notes that the existing debate about the appropriate use of millions of stored
10 specimens endures because of the uncertain and nature of past consents. Investigators and others
11 who collect and stored human biological materials now have the opportunity to correct past
12 inadequacies by obtaining more specific and clearly understood consents. By doing so, the need
13 to render samples unidentifiable may become less frequent, and the need to re consent minimized.
14 It is with these considerations that NBAC makes the following recommendations about
15 improving the consent process for the use of human biological materials in research.

16
17 **Recommendation 7:**

18
19 **To facilitate collection and storage of human biological materials in the future, consent**
20 **forms should be developed, as appropriate, to provide potential subjects with a**
21 **sufficient number of options to help them clearly understand the nature of the decision**
22 **they are about to make. Such options might include, for example:**

- 23
24 a) **To refuse use of their biological material in research;**
25 b) **To permit only unidentified or unlinked use of their biological material in**
26 **research;**

as valid for enrollment in research, but recognizes its moral value.

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- 1 c) **To permit coded or identified use of their biological material for one particular**
2 **study only, with no further contact permitted to ask for permission to do**
3 **further studies;**
- 4 d) **To permit coded or identified use of their biological material for one particular**
5 **study only, with further contact permitted to ask for permission to do further**
6 **studies; or**
- 7 e) **To provide prospective authorization for all future coded or identified use of their**
8 **biological material.**

9
10
11 This policy for existing samples should be supplemented with special attention to areas of
12 research considered sensitive or potentially objectionable to some. In other words, if the source
13 of an identifiable existing sample chose the option of not rendering the sample unidentifiable and
14 authorized future identifiable research uses, he or she would enjoy the additional protection
15 afforded by the requirement of specific consent for uses of the sample that might be considered
16 sensitive or objectionable. Such a category might include, for example, certain behavioral
17 genetics protocols, studies differentiating traits among ethnic or racial groups, or research on
18 stigmatizing characteristics such as addictive behavior.

19
20 Appropriate criteria should be used to determine whether re-contacting the individual is the
21 appropriate course of action. Additional concerns should be addressed when developing a plan
22 to recontact any individuals. For example, if explicit consent was never obtained for use of a
23 sample (because it met the requirements for waiver), IRBs should consider potential harms that
24 might arise should a subject learn, after the fact, that his or her material had been used in an
25 experiment, unbeknownst to him or her.

26

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1 ***Obtaining Consent in the Clinical Setting***

2 When samples are collected, whether in a research or clinical setting, it is appropriate to ask
3 subjects for their consent to future use of their sample, even in the case where such uses are at
4 the time unknown. The elements of the consent process for new samples should be the same as
5 those discussed previously for the use of existing identifiable samples.

6
7 There has been discussion in the literature and in testimony given before NBAC of the
8 concerns that arise when administering a consent process in a clinical setting (Transcripts Dec 9,
9 1997). These concerns often note that the clinical setting, where stress may be high, may not be
10 conducive to a consent process that involves complex choices about issues not directly related to
11 clinical care, and which involve thinking about the distant future. In this setting individuals may
12 be anxious about the clinical procedure and may not be prepared to consider carefully the factors
13 that go into making informed decisions about hypothetical research use of their tissue. The fact
14 that individuals will also be faced with other decisions and paperwork related to the clinical
15 procedure compounds the problem of administering an informed consent process in this setting.

16
17 Another way of improving the consent process may be to inform individuals about, and
18 ask for their consent to, future research use of their sample at some point before or after consent
19 is obtained for the clinical procedure. More studies should be done on the issue of the best time
20 to administer this consent in the clinical setting. NBAC acknowledges the important contribution
21 to this discussion of groups such as the National Action Plan for Breast Cancer, which has done
22 thoughtful work on ways to improve the overall consent process, including the timing of
23 obtaining consent. As investigators and IRBs consider this issue, it may be useful to consult the
24 work of groups who have made helpful suggestions regarding the design and timing of the
25 consent process. Using such guidance and their collective experience, the scientific community
26 should develop a consensus around a standard method for human biological material collection
27 in both therapeutic and research contexts that would minimize the need for complex recontact

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1 efforts.

2

3 **Recommendation 8:**

4

5 **When informed consent to the research use of human biological materials is**
6 **required, it should be obtained separately from informed consent to clinical**
7 **procedures.**

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1 **Recommendation 9:**

2
3 **When seeking informed consent in the clinical setting, it should be made clear to**
4 **subjects that refusal to consent to the research use of biological materials will in no**
5 **way affect the quality of their clinical care.**

6
7 **Criteria for Waiver of Consent**

8
9 When an investigator proposes to do research with identifiable samples, it is considered
10 research with human subjects. Ordinarily the potential research subject is asked whether he or
11 she agrees to participate. Seeking consent demonstrates respect for the person's entitlement to
12 choose whether to cooperate with the scientific enterprise, and it permits individuals to protect
13 themselves against unwanted or risky invasions of privacy. The adequacy of the requirement of
14 informed consent to provide appropriate protections should be evaluated in terms of whether or
15 not it achieves its intended goal. The purpose of informed consent in research is to provide
16 potential subjects with materially relevant information about the purpose and nature of a
17 proposed study, and appropriate information about risks and benefits to enable persons to make
18 a voluntary decision regarding participation. In considering the conditions for which informed
19 consent should be required for the research use of human biological materials, NBAC recognized
20 that informed consent, *by itself*, cannot provide protection for all the legitimate interests at stake
21 in the practice of gathering and using biological samples. Instead, informed consent plays an
22 important but not exclusive role in safeguarding both human subjects and research interests. Of
23 course, consent can never by itself protect someone from harm: it can only provide individuals
24 with available information about the probability and magnitude of harm. Overly elaborate
25 consent requirements cannot guard against all harms to subjects, would be extremely costly, and
26 could constrain socially valuable scientific research.

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1 As stated in the current federal regulations, human subjects research is presumed to
2 require consent, but this requirement can be altered or waived if all four criteria, set forth at 45
3 CFR 46.116(d), are met.

- 4
- 5 1) the research involves no more than minimal risk to the subjects;
 - 6 2) the waiver or alteration of consent will not adversely affect the rights and welfare of the
7 subjects;
 - 8 1) the research could not practicably be carried out without the waiver or alteration; and
 - 9 2) whenever appropriate, the subjects will be provided with additional pertinent information
10 after participation.

11

12 Determining the risks of research and the effects that waived consent might have on the
13 rights and welfare of the subject are bedrock considerations in deciding the level of protection
14 required for human subjects in research. Determining the level of risk to the subject is a key
15 criterion in deciding eligibility for expedited IRB review and in assessing the need to obtain
16 informed consent from the subject. Four key terms are central to this determination: “minimal
17 risk,” “rights and welfare,” “practicability,” and “after participation.”

18

19 ***Minimal Risk***

20 The regulations state that “*Minimal risk* means that the probability and magnitude of
21 harm or discomfort anticipated in the research are not greater in and of themselves than those
22 ordinarily encountered in daily life or during the performance of routine physical or
23 psychological exams or tests” (45 CFR 46.102(i)). Determining whether research risks are
24 minimal thus depends upon a comparison of research risks with risks which persons “ordinarily”
25 face outside of the research context.

26

27 However, when considering the risks of research conducted on human biological

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1 materials, one can question the applicability of the threshold that the regulations establish for
2 assessing minimal risk. The risks encountered “during the performance of routine physical or
3 psychological exams or tests” have limited utility as a baseline. While these risks can be
4 compared to the physical risks faced in the collection of new samples, they are not really
5 comparable with the risks of social and psychological harm relevant to research on biological
6 samples. The risks encountered “*during the performance*” of a medical exam evidently relate to
7 harms which the intervention itself may produce. The risks of psychosocial harm associated with
8 research on biological samples, on the other hand, relate to future uses of information derived
9 from samples.

10
11 The risks of “daily life” seem a more promising threshold for assessing the risks of
12 research on biological materials. In research on biological samples, the potential harms of central
13 concern (e.g., stigmatization, insurance and employment discrimination, familial conflict, anxiety,
14 violations of privacy) are those which can result if certain information from biological samples
15 (e.g., the subject’s susceptibility to disease) is disclosed to non-investigators. But such
16 information is also commonly contained in medical records. Persons (research subjects and non-
17 research subjects alike) generally face the risk that diagnostic, predictive, and other forms of
18 information about them contained in their medical records will be obtained and used in a harmful
19 manner. Although there are insufficient data to make a decisive statement about the relative
20 probabilities of harm resulting from uses of biological samples vis-a-vis access to medical
21 records, one might hold that the level of risk is similar in both cases. Indeed, research on
22 biological samples arguably poses lesser risks, since the sources of even “identifiable” samples
23 may be more difficult to trace than the subjects of explicitly labeled medical records. Thus, one
24 might conclude that most research on biological samples is “minimal risk.”

25
26 NBAC does not find this analysis of “minimal risk” to be compelling. On this reading of
27 the regulations, the issue is not fundamentally whether the risk of harm which research poses to

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1 subjects is in itself minor or substantial; rather, the issue is whether the risks the research presents
2 are more severe than risks which persons ordinarily confront outside of research. On this
3 interpretation, research risks could be substantial but nevertheless count as “minimal.” The
4 problem is that the purpose of assessing whether risk is “minimal” is to help IRBs determine
5 what types of protections should be required. While a strict reading of the regulations may permit
6 an interpretation which permits one to deem great risks of harm to subjects “minimal,” such an
7 interpretation certainly violates the spirit of the regulations.

8
9 An alternative interpretation of the regulations avoids this result. On this interpretation,
10 “‘risks of everyday life,’ has normative as well as descriptive force, reflecting a level of risk that is
11 not simply accepted but is deemed socially acceptable.”¹¹ According to this account any risk that
12 is not socially acceptable cannot properly be characterized as a risk of “daily life.” There is a
13 widespread view that the present risks of harm from uses of sensitive medical information about
14 individuals are not acceptable, and that we need stronger privacy laws to remedy this situation.
15 Thus, the risks of harm resulting from the improper use of medical records are not, on this
16 interpretation, risks of “daily life.” It follows that one cannot employ the risks of harmful uses of
17 medical records as a baseline for determining whether research on biological samples is minimal
18 risk. This, in turn, makes it difficult to perform a minimal risk analysis for research on biological
19 samples, as there are no apparent alternative candidates that can plausibly serve as a baseline.

20
21 **Recommendation 10:**

22
23 **Institutional Review Boards should, in general, operate on the presumption that**
24 **research on *existing* coded samples is of minimal risk to the human subjects if: 1)**
25 **the study adequately protects the confidentiality of personally identifiable**
26 **information obtained in the course of research; 2) the study does not involve the**

¹¹ Benjamin Freedman, Abraham Fuks, Charles Weijer, “In loco parentis: Minimal Risk as an Ethical Threshold for

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1 **release of information to any third party with an interest in the employment or**
2 **insurability of the human subject; and 3) the study design incorporates an**
3 **appropriate plan for whether and how to reveal findings to the sources or their**
4 **physicians should the findings merit such disclosure.**

5
6 While the regulatory definition of “minimal risk” thus appears inadequate for research on
7 human biological materials, the additional requirement that the waiver of consent must “not
8 adversely affect the rights and welfare of the subjects” (45 CFR 46.116 (2)(d)(2)) is sufficient to
9 protect the same interests. As discussed below, the rights and welfare condition for waiver or
10 alteration of consent requires an assessment of the risks of psychosocial harms and protects
11 subjects from any substantial risks.

12 13 ***Rights and Welfare***

14 Failing to obtain consent may adversely affects the rights and welfare of subjects in two
15 basic ways: (1) The subject may be improperly denied the opportunity to choose whether to
16 assume the risks that the research presents; (2) The subject may be harmed or wronged as a result
17 of their involvement in research to which he or she has not consented.

18
19 A waiver of consent in the collection of *new* biological samples violates subjects’ rights
20 because it would expose them to unwanted bodily invasions. The interest in being free from
21 unwanted bodily invasions is the primary interest the requirement of informed consent was
22 instituted to protect. In the case of consent for the use of *existing* samples, the interests at stake
23 are different. In this context, it is principally the social and psychological harms delineated in
24 Chapter 4 that are at issue. Subjects’ interest in controlling information about them is tied to their
25 interest in, for example, not being stigmatized or not being discriminated against in employment
26 and insurance. The degree to which the assertion of these interests is compelling is a function of

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1 the probability of harm occurring. Important considerations that figure into the probability of
2 harm occurring, include:

3
4 (1) How easily is the sample source identifiable?

5 (2) What is the likelihood that the sample source will be traced?

6 (3) If the source is traced, what is the likelihood that persons other than the investigators
7 will obtain information about the source? (Privacy/confidentiality laws may be relevant
8 here, as is the integrity of investigators and their institutional confidentiality protections.)

9 (4) If non-investigators obtain the information about the source, what is the likelihood that
10 harms will result, including adverse consequences arising from the reporting of uncertain
11 or ambiguous clinical results? (State and federal discrimination laws may be relevant with
12 respect to uses of information by third parties).

13
14 As noted in Chapter 4, the probability of psychosocial harms resulting from research on
15 biological samples is somewhat speculative at present. There are, however, good reasons to think
16 that the risks of harm are generally minimal, or at least can easily be rendered minimal. Given
17 current scientific practices, there are few studies where it is necessary that investigators know the
18 identity of sample sources. Thus, investigators will not usually have a need to trace sample
19 sources although they might require additional clinical information without identifying the
20 source. Even where investigators do trace a source, it is not necessary to reveal information
21 about sources to third parties. While it is nonetheless possible that non-investigators will access
22 information about a source, investigators can minimize this risk through appropriate
23 confidentiality mechanisms. For example, protocols that include provision for a way to isolate the
24 results of genetic or other research results completely from the subject's medical record, and that
25 incorporate a prohibition on returning uncertain or ambiguous information to subjects (which
26 would forestall the communication of premature and potentially upsetting information) should in
27 most cases ensure that risks will be minimal.

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Although the risks of psychosocial harms may generally be minor in research on human biological materials, there are some important exceptional cases. For example, controversial studies such as those which involve behavioral genetics or which make explicit comparisons between ethnic or racial groups, are likely to offend some research subjects and threaten their ascriptive identity. Moreover, there remains the likelihood that the results of such studies will be used to stigmatize and discriminate against group members (research subjects and non-research subjects alike).

Recommendation 11:

In considering waiver of consent, the term “adversely affects the rights and welfare” of human subjects should be interpreted to mean that the waiver does not violate any state or federal statute or customary practice regarding entitlement to privacy. Considerations of rights and welfare should also include an assessment of the potential effects of a study that examines traits commonly considered to have political, cultural, or economic significance to the community to which the sample source belongs.

Practicability

An investigator who requests a waiver of the informed consent requirement for research use of human biological materials under the current federal regulations must provide to the IRB evidence that it is not practicable to obtain consent. Neither the regulations nor OPRR offer any guidance on what defines practicability.¹²

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1 Practicable is defined in the ordinary sense as that which “can be done or used,” or is
2 “possible in practice” (*Oxford English Reference Dictionary*). This could suggest that obtaining
3 consent is always practicable, so long as there are the means and skills to carry this out, but that it
4 can never be an absolute requirement. The issue for regulatory purposes, and, NBAC would
5 suggest, for the purpose of assessing the ethical acceptability of this provision, is whether the
6 practicability requirement—alone or in combination with other criteria for obtaining a waiver—
7 adds guidance to the investigators and IRBs who will make these decisions. Informed consent
8 may not be “possible in practice” when there are many more subjects than there are individuals
9 to seek their consent, or when the amount of time it would take to recontact would be longer than
10 the period of time the study was to take place. Similarly, obtaining consent might be thought of
11 as impracticable if the financial costs either of a direct recontact effort, or even indirect efforts
12 (such as mailing consent forms and information) far exceeded the researcher’s budget. One
13 might even suggest that in research that is designed to hold out the prospect of direct benefit to
14 some of the subjects, it would be impracticable to take the time to recontact potential subjects
15 because the delay in completing the study could be thought of as a more serious harm than the
16 failure to obtain express consent. While these are reasonable examples of impracticability, and,
17 NBAC would suspect, might be regarded by some as good reasons for granting a waiver, the
18 trouble with the practicability requirement is that it forces a comparison between otherwise
19 incommensurable harms: the wrong that could be committed by not obtaining informed consent,
20 and the prohibitively costly, perhaps difficult, and even needlessly intrusive harm of attempting
21 recontact. As with many types of incommensurability in IRB review the customary task of
22 assessing risk and benefit becomes far more problematic.

23

24 **Recommendation 12:**

25

26 **If research using *identifiable, existing* human biological materials is determined to**

¹². Personal Communication, Dr. Gary Ellis, Director, OPRR, August 25, 1998.

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1 **present minimal risk, Institutional Review Boards may presume that it would be**
2 **impracticable to meet the consent requirement (45 CFR 46.116(d)(3)). This**
3 **interpretation of the regulations applies only to the use of human biological materials**
4 **collected before the adoption of the recommendations contained in this report**
5 **(specifically Recommendations xx regarding informed consent). Materials collected**
6 **after that point must be obtained according to the recommended informed consent**
7 **process and, therefore, Institutional Review Boards should apply their usual**
8 **standards for the practicability requirement**
9

10 Even where it might be deemed practicable to obtain consent for research use of stored
11 human biological materials, it may be unnecessarily burdensome for investigators. NBAC
12 believes that in assessing the appropriateness of waiving consent, consideration should be given
13 principally to the criteria of minimal risks and rights and welfare. Practicability should not be a
14 compelling consideration.
15

16 NBAC recognizes that if its recommendation that coded samples are identifiable is
17 adopted, there may be an increase in the number of protocols that require IRB review. If,
18 however, such a protocol is then determined by the IRB to present minimal risk to a subject's
19 rights and welfare, the requirement for consent may be waived if the practicability requirement is
20 revised for this category of research.
21

22 NBAC believes that these interpretations and recommended changes in the regulations
23 will allow important research to go forward while still taking into consideration potential harms to
24 subjects. However, it must be noted that by dropping the requirement that consent must be
25 obtained if practicable, NBAC does so with the expectation that the process and content of
26 informed consent for new studies will be explicit as to the intentions of the subjects regarding the
27 research use of their samples (see Recommendations concerning consent).

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Providing Additional Information as Required at 45 CFR 46.116(d)(4)

In the current regulations, the third condition for the waiver of consent stipulates that, “whenever appropriate, the subjects will be provided with additional pertinent information after participation.” The historical context for this condition are “deception” studies (e.g., the behavioral sciences) in which it is deemed crucial to study design that the individual not know of their status as a research subject. Thus, according to the regulations, the IRB, while waiving consent (by finding and documenting the first three required conditions), could require that subjects be informed that they were subjects of research, a so-called “debriefing” requirement.

The applicability of this condition in the context of stored samples could be interpreted in a variety of ways. If the first three conditions of waiver of consent are met, the IRB might require, as an additional measure of protection, that the investigator provide further information to the subjects. Such a communication would describe the status of the research project and inform them that their samples will be used or were used in the research. Such a requirement might only be appropriate if consent had already been obtained and the IRB determines that re-consent is not required for a specific or new protocol. The IRB might well recognize that only those subjects who could be found would be so informed.

Recommendation 13:

The Office for Protection from Research Risks should make clear to investigators and Institutional Review Boards that the fourth criterion for waiver, that “whenever appropriate, the subjects will be provided with additional pertinent information after participation,” (45 CFR 46.116(d)(4)) usually does not apply to research using human biological materials.

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1 This criterion was designed to address the situation in which an IRB has permitted
2 informed consent to be waived or modified with the result that subjects may be aware of
3 interacting with an investigator but still deceived about the true nature of the research, or in which
4 subjects will be unaware of being observed in public settings. (check history) Respect for
5 subjects' rights and welfare in such circumstances will usually dictate that they be informed after-
6 the-fact of the research in which they have been involved as "naive" or unwitting subjects, and
7 perhaps offered the opportunity to withdraw their information from the investigator's data. In
8 general, however, NBAC concludes that this fourth criterion for waiver on consent is not relevant
9 to research using human biological materials, and, in fact, might be harmful if it forced
10 investigators to recontact individuals who might not have been aware that their materials were
11 being used in research.

12

“Opt Out” as an Additional Measure of Protection when the Consent Requirement Has Been Waived

15 “Opt out” refers to the choice given to subjects to exclude themselves from a study.
16 Unless someone has “opted out,” he or she is assumed to be enrolled. If, after a waiver of the
17 consent requirement is granted, an investigator or IRB has residual concerns about the nature of
18 the research or the possibility that some individuals might find the research objectionable, then an
19 additional measure can be taken to allow subjects to opt out of the research. In this scenario,
20 subjects would, if possible, be contacted and given the choice of opting out; if they did not
21 respond or could not be found, the sample could still be used because the consent requirement
22 had already been waived. This differs significantly from a scenario in which the consent
23 requirement has not been waived. In that scenario, if a person did not respond with explicit
24 consent or could not be found, his or her sample could not be used in the research protocol.

Rendering Existing Identifiable Samples Unidentifiable to Avoid the Need for Consent

26 A more practical solution to using existing samples for which it is impracticable or

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1 problematic to gain express informed consent for a specific use of the sample is to render the
2 samples unidentifiable. The rationale for this apparently simple proposal is that in many cases
3 existing samples were collected without anything resembling adequate disclosure that they would
4 be used for a range of purposes unrelated to the context in which they were collected.

5
6 There are several drawbacks to rendering existing samples unidentifiable for every use
7 that is not specifically consented to by the source. First, there is the administrative cost of
8 rendering such samples completely unidentifiable. Second, if a sample is not identifiable,
9 opportunities may be lost to protect the well being of the source or his or her relatives (e.g., in the
10 case of genetic conditions) when later research discovers therapeutically significant links between
11 various diseases or between diseases and genotypes. Third, rendering a sample unidentifiable
12 restricts the usefulness of that sample to investigators, who might wish to obtain additional
13 samples, or who might wish to gather additional medical information from the patient or the
14 medical record. Thus, there could be a scientific or medical price to pay for this action. Fourth,
15 some investigators may choose to render identifiable samples unidentifiable so as to avoid the
16 time and cost of IRB review and the possibility that the IRB may require obtaining informed
17 consent.

18
19 Another possible ethical objection to this practice is based on the belief that rendering
20 existing samples unidentifiable without consent is problematic because researchers once had the
21 opportunity to seek consent but did not exercise it.

22
23 NBAC believes that rendering existing samples unidentifiable in order to expedite
24 research protocols can be avoided in many situations by designing the research in such a way as
25 to minimize risks to the subjects. If risks are minimal, then it is possible that the requirement for
26 informed consent might be waived or altered according to the regulations, 45 CFR 46.116(d). If
27 the nature of the research changes in the future, so that an investigator now selects specific

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1 samples for additional studies that might increase risks beyond the minimal level, further IRB
2 review would be required.

3
4 Moreover, for future sample collection, a consent process that is explicit about the
5 subject's wishes concerning permissible uses of tissue will help to alleviate the need for the
6 investigator to use unidentifiable samples.

7
8 Nevertheless, the NBAC recognizes that there will be some situations in which it is
9 scientifically sound or desirable to render samples unidentifiable, and there is no scientific or
10 medical cost to doing so. In addition, NBAC recognizes that going back to seek consent could be
11 costly and time consuming in situations where there is a small possibility for stigmatization or
12 harm once the identifiers are removed. Furthermore, contacting individuals might be disruptive
13 and even unwanted by the sample source. With these considerations in mind, NBAC concludes
14 that, in those circumstances where valuable samples could not otherwise be used, where consent
15 would be difficult to obtain, and where there is no scientific cost to losing the link, it is ethically
16 acceptable to render samples unidentifiable without the source's consent. In arriving at this
17 conclusion, NBAC also considered public input it received during deliberations, in which most
18 people emphasized that they did not view their donated biological material as something that
19 belonged to them, but rather as a gift to be used by the scientific community subject to the review
20 for quality and ethical acceptability, and if they could be assured that the information obtained
21 would not be used to discriminate against them.

22
23 **Recommendation 14:**

24
25 **When samples are to be drawn from identifiable specimens, investigators who**
26 **choose to have the identifiers stripped from the samples should explain to the**
27 **Institutional Review Board the decision not to work with the samples on a coded or**

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1 **identified basis.**

2 **Reporting Research Results to Subjects**

3

4 Experts disagree about whether interim or clinically inconclusive findings from research
5 should be communicated to subjects, although most agree that such findings should not be
6 conveyed because only confirmed, reliable findings constitute clinically significant or
7 scientifically relevant information. Persons who oppose revealing interim findings argue that the
8 harms that could result from revealing preliminary data are serious, including anxiety or
9 unnecessary (and possibly harmful) medical interventions. They prefer to avoid such harms by
10 controlling the flow of information to subjects and limiting communications to those that
11 constitute reliable information. MacKay (1984), writing about the development of genetic tests,
12 argues against revealing interim findings, contending that preliminary results do not yet constitute
13 “information” since “until an initial finding is confirmed, there is no reliable information” to
14 communicate to subjects, and that “even...confirmed findings may have some unforeseen
15 limitations” [p. 3]. Subjects should not be given information about their individual test results
16 until the findings have been confirmed through the “development of a reliable, accurate, safe and
17 valid presymptomatic test” [pp. 2-3; see also Fost and Farrell (1990)]. Others have argued that the
18 principle of autonomy dictates that subjects have a right to know what has been learned about
19 them, and therefore, that interim results should be shared with subjects (Veatch).

20

21 Reilly (1980) suggests that IRBs develop general policies governing the disclosure of
22 information to subjects to help make these determinations. At least the following three factors
23 should be considered: “1) the magnitude of the threat posed to the subject; 2) the accuracy with
24 which the data predict that the threat will be realized; and 3) the possibility that action can be
25 taken to avoid or ameliorate the potential injury” [p. 5]. IRBs should ask investigators to define
26 three categories of findings: 1) “findings that are of such potential importance to the subject that

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1 they must be disclosed immediately;” 2) “data that are of importance to subjects..., but about
2 which [the investigator] should exercise judgment about the decision to disclose....[i]n effect,
3 these are data that trigger a duty to consider the question of disclosure;” and 3) “data that do not
4 require special disclosure” [pp. 5, 12].

5
6 **Recommendation 15:**

7
8 **Institutional Review Boards should develop general guidelines for the disclosure of**
9 **the results of research to subjects and require investigators to address these issues**
10 **explicitly in their research plans. In general, these guidelines should reflect the**
11 **presumption that the disclosure of research results to subjects represents an**
12 **exceptional circumstance. Such disclosure should occur only when all of the**
13 **following obtain:**

- 14 **a) the validity and clinical significance is high;**
15 **b) the threat to the subject’s health, as indicated by the research finding, is**
16 **significant; and**
17 **c) there is readily available a course of action to prevent, avoid, ameliorate, or treat**
18 **the threat to the subject’s health.**

19
20 **Recommendation 16:**

21
22 **The research protocol should describe anticipated research findings and**
23 **circumstances that might lead to a decision to disclose the findings to a subject, as**
24 **well as a plan for how to manage such a disclosure.**

25
26 **Recommendation 17:**

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1 **When appropriate, persons should be asked whether they would be interested in**
2 **receiving research results if such disclosure is deemed appropriate by the**
3 **investigator.**

4

5 **Recommendation 18:**

6

7 **When research results are disclosed to a subject, appropriate medical advice or**
8 **referral should be provided.**

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1 Considerations of Potential Harms to Others

2
3 The federal regulations governing the protection of research subjects extend only to
4 individuals who can be identified as the source of the biological samples. The exclusive focus of
5 the regulations on the individual research subject is arbitrary from an ethical standpoint, since
6 persons other than the subject can both benefit and be harmed as a consequence of the research.

8 *Risks to Groups*

9 Research on samples that implicate groups may place group members at risk of harm.
10 For example, research revealing that a racial or ethnic group is unusually prone to disease could
11 be used to stigmatize and discriminate against group members.

12
13 OPRR guidance to IRBs and investigators on how best to identify and minimize risks to
14 groups is required. Consultation with group members prior to designing and implementing
15 research on groups, for example, may often be an effective way to understand and reduce risks to
16 groups. However, work needs to be done to identify appropriate mechanisms for group
17 consultation.

18
19 It also seems appropriate to highlight how some of these issues ought to be discussed
20 among researchers and their professional organizations. For example, what is the appropriate
21 role of public health policy in developing new knowledge from genetic epidemiology? Will
22 additional ethical considerations be adjusted to ensure that the benefits of public health objectives
23 do not come at the cost of individual concerns? For many studies, the answer may be yes: the
24 net gain to a particular “population” from knowing about its increased risk (especially when
25 something can be done at an individual level with this knowledge) will often outweigh the harms
26 that come from labeling a group as “high risk.”

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Risks and Potential Benefits to Relatives of the Sample Source

Others who may be at some risk are first-degree relatives, or next-of-kin. The need to consider these people “at risk” is particularly evident when the disease or condition being studied is genetic (and thus may be shared by family members) or diseases that involve infectious agents or toxic exposures. In these instances, investigators are likely to be fully aware that the research they are conducting on a sample might have implications for those closely related to the sample source, individuals who are readily identifiable.¹³ NBAC does not assume that because there might be risks to relatives of the sample source, those risks warrant considering those individuals to be human subjects, deserving the protection of informed consent.¹⁴ In fact, NBAC finds the possibility that a relative of the sample source could stop a research protocol on the basis of consent not only impractical, but also troublesome. If the sample source has consented to the research use of his or her sample, that consent alone is sufficient for the research to proceed. However, although the regulations do not require that the concerns of first-degree relatives to be considered, NBAC recognizes that there might be circumstances in which an investigator finds it useful, beneficial, appropriate, and feasible to consider potential harms and benefits with such individuals.

A different set of concerns arise when the source of the sample is deceased. Under the federal regulations, people are human subjects only while living. Research involving human biological materials from individuals who are deceased at the time of the research is not subject to the requirements of DHHS regulations, regardless of whether or not prior informed consent was

¹³ This distinction is worth noting. In the case of membership in a group, persons might not be individually identifiable although identified as a member of that group. In the case of biological relatives, persons related to the sample source are likely to be individually identifiable. See DeRenzo, E.G., Biesecker, L.G., and N. Meltzer, “Genetics and the Dead: Implications for Genetics Research with Samples from Deceased Persons,” *American Journal of Medical Genetics* 69:332-334, 1997

¹⁴ OPRR has indicated that the living relatives might in fact be considered human subjects by virtue of their genetic relationship to the sample source, but the regulations—specifically the *OPRR Institutional Review Guidebook* section on human genetic research (pp. 5-42 to 5-63)—do not clearly specify how this consideration is to be handled by IRBs.

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1 obtained. In addition, the existing regulations do not make explicit the status of living relatives of
2 deceased individuals whose stored samples are used in research.¹⁵ However, it is possible that the
3 living relatives of the deceased sample source might have an interest in the research, particularly if
4 the investigation focused on hereditary traits.

5
6 **Recommendation 19:**

7
8 **Research using coded and identified samples, even when not potentially harmful to**
9 **individuals from whom the samples are taken, may be potentially harmful to groups**
10 **associated with the individual. Likewise, although individuals from whom unidentified**
11 **or unlinked samples are obtained are very unlikely to be harmed by research, studies**
12 **using such samples are potentially harmful to their associated groups. To the extent**
13 **possible, investigators should plan their research so as to minimize such harm and**
14 **should consult, when appropriate, representatives of the relevant groups regarding**
15 **study design.**

16
17 **Recommendation 20:**

18
19 **If it is anticipated that a specific research protocol poses a risk to a specific group, this**
20 **risk should be disclosed during any required consent process.**

21
22 **Publication and Dissemination of Research Results**

23
24 Publishing identifiable information in biomedical journals may pose a risk to the privacy and
25 confidentiality of research subjects. Publicly disclosing such information through written description

¹⁵ Please note 45 CFR 46.102 “Definitions: (f) Human subject means a living individual about whom an investigator (whether professional or student) conducting research obtains (1) data through intervention or interaction with the individual, or (2) identifiable private information . . .” (OPRR Reports, Protection of Human Subjects, 1991).

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1 or pedigrees may result in adverse psychosocial effects and, without informed consent of the
2 individual to do so, infringes on the rights of the subject or patient (Botkin et al). Because of the
3 familial nature of information in pedigrees, their publication poses particularly difficult questions
4 regarding consent. Journal editors have an ethical obligation to publish only that human subjects
5 research that they have reason to believe was conducted according to ethical standards set forth in
6 the Common Rule, which includes review by an Institutional Review Board (IRB). Recent studies
7 have reported that ethical standards communicated in journals' instructions to authors vary widely,
8 as does how well authors adhere to such standards.

9
10 **Recommendation 21:**

11
12 **Plans for disseminating results of research on human biological materials should**
13 **include, when appropriate, provisions to minimize the potential harms to individuals**
14 **or associated groups.**

15
16 **Recommendation 22:**

17
18 **When accepting research results for publication, journal editors should require**
19 **investigators to indicate whether the research was conducted in compliance with the**
20 **substantive requirements of the Federal Policy for the Protection of Human Subjects**
21 **in Research, even if the study was privately funded and exempt from the Federal**
22 **requirements for that research.**

23
24 **Professional Education and Responsibilities**

25
26 Public and professional education is an essential part of effective public policy on the use
27 of human biological materials for research. By education, NBAC is referring not simply to the

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1 provision of information with the aim of adding to the net store of knowledge by any one person,
2 or group; rather, education refers to the ongoing effort to inform, challenge, and engage.
3 Education about ethical issues in research involving human biological materials means that a
4 variety of individuals and groups would have new tools to assess these important issues.
5 Therefore, opportunities for such education need to be directed to IRBs, researchers, other
6 members of the research and academic community, political decision makers at the state and
7 federal levels, interest groups, possible human subjects and the eventual consumers of research
8 on human biological materials. There must be widespread and continuing deliberation and the
9 provision of information and education to the public in the area of genetics, and on other
10 developments in the biomedical sciences, especially where these affect important cultural
11 practices, values, and beliefs.

12

13 **Recommendation 23:**

14

15 **The National Institutes of Health , professional societies, and health care**
16 **organizations should continue and expand their efforts to train investigators about**
17 **the ethical issues and regulations regarding research on human biological materials,**
18 **and to develop exemplary practices for resolving such issues.**

19

20 NIH can promote these efforts through the use of such mechanisms as workshops,
21 requirements for training grants and center grants, and funding for research on pertinent topics
22 related to this report. Professional societies can develop training materials on these issues and
23 disseminate information about how research centers have successfully addressed ethical issues
24 regarding research on human biological materials. Special emphasis should be given on
25 developing consent processes that allow patients and research volunteers to make meaningful
26 choices about how biological materials might be used in future research. Continued collaborative
27 efforts between scientists and patient representatives and advocacy groups are likely to be

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1 particularly fruitful in strengthening the consent process.

2
3 These discussion should encompass the kinds of issues raised by storage and use of
4 human biological materials and the implications of such research on important values. Moreover,
5 as it is the research community that seeks access to these materials, for policy purposes a moral
6 burden should fall on researchers to elicit from prospective contributors, both individual and
7 communal, the values and meaning they attach to the requested samples.

8
9 **Recommendation 24:**

10
11 **Compliance with the recommendations set forth in this report will require additional**
12 **resources. All research sponsors (government, private sector enterprises, and**
13 **academic institutions) should work together to make these resources available.**

14
15 **Use of Medical Records in Research on Human Biological Materials**

16 {Language to be written after text in chapter 3 is updated and revised.}

17
18 **Recommendation 25:**

19
20 **Because research using identifiable human biological materials sometimes requires**
21 **that investigators have access to information in a patient's medical record, state and**
22 **federal legislation concerning medical record privacy should include provisions for**
23 **legitimate access by researchers who have met all applicable review and consent**
24 **requirements.**

25
26 **Recommendation 26:**

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1 **State and federal legislators are encouraged to enact statutes on medical records**
2 **research and human biological materials research that are uniform in their**
3 **approach, while taking into account that the 'information' that can be found by**
4 **studying cells and tissues differs in important respects from that contained in**
5 **medical records.**

7 **Conclusions**

8
9 To advance human health it is critically important that human biological materials
10 continue to be available to the biomedical research community. It increasingly will be essential
11 for investigators to collect human biological materials from individuals who are also willing to
12 share important clinical information about themselves. In addition, it is crucial that the more than
13 282 million samples already in storage remain accessible under appropriate conditions.

14
15 The growing availability to third parties of genetic and other medical information about
16 individuals has fueled the current debate about medical privacy and discrimination. As a society
17 we are sensitive to the possibility that the use of information obtained from human biological
18 samples can lead to harms as well as benefits. These concerns require that those who agree to
19 provide their DNA, cells, tissues, or organs for research purposes not be placed at unacceptable
20 risk. Measures to provide appropriate protections for individual privacy and for the
21 confidentiality of clinical and research data are important if significant research is to continue.
22 The recommendations provided in this report are intended to promote the goals of improving
23 health through biomedical research while protecting the rights and welfare of those individuals
24 who contribute to human knowledge through the gift of their biological materials.