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Sharing Individual Research Results with Biospecimen Contributors: Point

Rihab Yassin¹, Carol Weil², and Nicole Lockhart²

Biospecimens are an essential resource for biomedical research, including research that aims to decipher the biology of cancer and improve its clinical management. Accordingly, patients with cancer and others who contribute biospecimens for research are increasingly attentive to the results of studies conducted with their biologic samples (1). They may seek aggregate research results as well as individual results, including incidental findings uncovered in the course of research (2–4). As advanced technologies for cancer research make more feasible the discovery of biomarkers for risk assessment, diagnosis, and prognosis, the research community is faced with the challenge of whether to share specific research results with biospecimen contributors.

Sharing research results, particularly genetics and genomics, with research participants has been at the forefront of many ethical and scientific discussions. Advocates of the process maintain that research participants should be offered the option of receiving potentially important information that could impact their health and life choices (1, 5–7). Moreover, investigators often feel motivated to share results that they perceive to be beneficial to research participants not only because of their potential clinical value but also to promote a trusting partnership with them. Opponents of sharing results assert that the purpose of research is to generate knowledge rather than provide clinical care and that research laboratories do not necessarily operate in accordance with clinical laboratory standards (1). In the United States, the Clinical Laboratory Improvement Amendments (CLIA) law (8) and implementing regulations (9) require a “laboratory” as defined in these authorities to obtain federal certification and accreditation before disclosing results for patient care. However, the legal definition of “laboratory” in the CLIA statute and implementing regulations limits this mandate to facilities that analyze human tissue for the purpose of clinical diagnosis, prevention, or treatment of disease, or impairment or assessment of health (8). Arguably, many research biobanks would not fit this definition as they do not analyze tissue for the purpose of clinical care. Furthermore, facilities only collecting or preparing specimens and not conducting testing are expressly exempt from CLIA requirements (9). Accordingly, we postulate that CLIA may not require biobanks or research facilities to be CLIA-certified to share research results with participants, as long as such entities are not expressly constituted for the purpose of conducting clinical laboratory testing. While there is controversy about the legal question of whether non–CLIA-certified facilities may return research results, it has been recognized that results from non–CLIA-certified laboratories can still have analytic validity and that it may be appropriate as an ethical and policy matter for biobanks and studies to return such results labeled as “research” (4, 6).

Even for CLIA-certified laboratories, there is disagreement about the nature and appropriate scope of research results that can be shared with participants. Some clinical results can be irrefutably beneficial from the standpoint of clinical use—for example, informing research participants about a cancer predisposing genetic mutation with available preventive measures that can avert fatal disease. There is, however, less consensus about the return of results with unclear medical significance or where clinical intervention is unavailable. For example, sharing research results about genetic predisposition to early onset Alzheimer disease might permit lifestyle adjustments and financial planning in anticipation of disease progression but is not clinically actionable. These complex issues underlie the importance of developing overarching principles for the sharing of research results to guide the scientific community.

We advocate that attention be given to the conception of best practices for sharing research results during the planning and development of biobanks and for any research projects that involve the collection of biospecimens. These best practices should consider both scientific and ethical elements. Scientific elements include the objective of the research project, the type and identifiability of the biospecimens used, whether biospecimens are collected for specific or unspecified future research, the type of data generated (i.e., have clinical/health relevance), and the scientific validity and reproducibility of the data. Ethical elements include the nature of the informed consent process for the collection of biospecimens and the extent to which it offers clear options to research participants for receiving, declining, or being re-contacted about the receipt of research results. In addition, the ethical mandate to protect the welfare of human subjects

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(10) requires consideration of whether sharing research results will cause unwarranted assurance or unnecessary alarm.

To address some of these tangible issues, the National Cancer Institute (NCI) organized a workshop in 2010 entitled “Release of Research Results to Participants in Biospecimen Studies” with the objective of developing best practices about when and how to share biospecimen research findings with study participants. The workshop included a discussion group devoted to the conditions for sharing individual research results from biospecimens collected for/in the course of clinical studies (11). Herein, we describe a framework for decision making about whether and when to return research results from identifiable biospecimens that is based on recommendations from the NCI workshop. This framework incorporates a set of operational criteria comprising a “smart filter” to guide the sharing of individual research results from biospecimens collected for/in the course of clinical studies (11). These criteria include the following 4 core elements: (i) the research participant agreed in the consent process to receive research results; (ii) the result is analytically valid; (iii) the result is clinically significant or serious for the participant; and (iv) the result is clinically actionable (11).

With regard to the first core element, the informed consent form should describe if research participants may choose to receive, or be re-contacted about, research results. The consent form is internationally recognized as the operational mechanism for ensuring informed and voluntary decisions to participate in research (12, 13). The NCI workshop recommended that the informed consent document contain “language to describe the circumstances under which research participants may receive individual, clinically relevant research findings and the mechanisms for communicating such findings” (11). Yet, particularly in genome-wide studies and research analyzing genetic databases in conjunction with medical records, there are inherent limitations to disclosing the full extent and nature of possible research findings. Considering the current suggestion of the HHS Office for Human Research Protections that one-time general consent for future research use of biospecimens be required (14), the universe of potential research results might be virtually indescribable. Moreover, increasingly sophisticated analytic platforms and informatics tools have transformed cancer research, complicating the common understanding of typical consent form descriptors such as “genetic research.” There is a great need for research to determine how participants interpret such consent form language. Given the ambiguities inherent in communicating the intention to share results, when a specific finding falls within the scope of the informed consent, participants should receive an additional opportunity to consider whether to receive or decline known information, particularly unexpected incidental findings. Investigators collecting biospecimens should consider during the protocol design stage whether it would be ethically appropriate to offer research participants the option to decline the receipt of research results with potential clinical use, such as diagnostic discrepancies. Some have argued that in rare cases, a “duty to rescue” exists requiring the return of

![Figure 1](https://www.aacrjournals.org)
research results that offer clear evidence of clinical use and actionability (15).

With regard to the second core element, ensuring the analytic validity of research results, standards may not exist in every research laboratory for assessing the accuracy, precision, sensitivity, and specificity of the testing conducted and results generated. When research results are generated and interpreted in a non–CLIA-approved laboratory, the NCI workshop recommended that such results be verified in a CLIA-approved laboratory unless the verification is not possible. In such cases, investigators should ascertain the analytic validity and safety of the performed test (11). At a minimum, any research results generated in a non–CLIA-approved laboratory should clearly be labeled as research and not clinical findings (4, 6). An issue to consider in the context of CLIA verification is who will cover the cost of re-testing.

The third core element requires an assessment that the research result is clinically significant or serious. Interpreting the parameters of this element can be highly subjective and therefore should be undertaken in consultation with an Institutional Review Board (IRB) or other expert panel. Factors to consider include the relative risk of the disease or health condition associated with the information, the likelihood of fatality, and the level of morbidity. The NCI workshop did not reach consensus about whether to share results that are not clinically significant to the research participant but have reproductive implications or significance to family members. Individual and societal values of the research participant and any financial constraints should be considered by biospecimen projects in deciding whether to broaden the scope of sharing to include these criteria.

The fourth core element restricts the sharing of individual research results to those that are “clinically actionable.” As with the third element, defining the parameters is subjective, and no clear consensus was reached by the NCI workshop. A more narrow view restricts returnable results to those that lead to therapeutic interventions designed to cure disease. This is consistent with a perceived duty of medical researchers to deliver information that clearly shows the benefit of available remedies (15). A broader view would consider appropriate the sharing of information which, even if it would not support specific therapies, would help research participants manage their medical condition or plan for the future. Sharing the greater range of information would be more appropriate for biospecimen research projects that can provide access to genetic counselors and medical staff, or otherwise assist research participants with the process of interpreting and understanding research results.

Every project faces unique issues in determining how, when, and by whom research results can best be shared. Biobanks housed in university hospitals, cancer centers, and community hospitals, and those specially created for multicenter clinical trials, rare disease research collaborations, or other specific studies, will differ significantly in infrastructure, scope of mission, and funding sources. Whenever possible, research results should be communicated by knowledgeable professionals who have ongoing relationships with participants. Such individuals are also most likely to be in possession of private identifiable information about participants and therefore will be in a position to contact participants with fewer legal constraints. Policies for sharing individual research results should be developed in consultation with local IRBs, advisory boards with representatives from the participant community or other representative groups.

In terms of who is responsible for communicating research results to participants, the NCI workshop recommended that the biospecimen collection sites be responsible for communicating results generated from primary research or from an internal review of biospecimens (e.g., pathology) at the biobank (11). In general, results can be linked to individual participants by the primary collecting site that maintains identifiable information or in some cases by the biobank, if it holds identifiable information. The NCI workshop also considered whether biobanks should assume the responsibility for sharing results generated in the course of secondary research studies. Workshop attendees recommended that biobanks coordinate the process by which secondary research results are communicated to human research participants with input from the primary and secondary institutions (11). In most cases, secondary researchers would not be able to link research results to individual participants as they would likely receive coded or de-identified samples. The NCI workshop discussed mechanisms such as the material transfer agreement as a means of clarifying the responsibilities of all participants in the research process (11).

In conclusion, limiting the scope of research result sharing to defined criteria such as those described in the “smart filter” shows respect for research participants and protects them from needless harm that can be caused by preliminary or nonvalidated research findings. The recommendations generated by the NCI workshop (11) will be incorporated in a future iteration of the NCI Best Practices for Biospecimen Resources (16) to guide the scientific community. More research is needed to further the development of evidence-based criteria for research results that can be shared with research participants.

Disclosure of Potential Conflicts of Interest

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9. CLIA regulations, 42 CFR 493.2.