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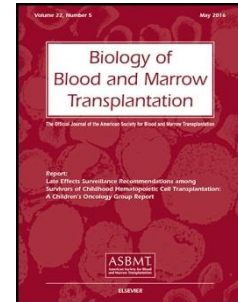
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Improving the Informed Consent Process in Hematopoietic Cell Transplantation: Patient, Caregiver, and Provider Perspectives

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Running Title: Patient, Caregiver, and Provider Perspectives on Informed Consent

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Highlights

- Important information to clinical research participants includes benefits to others
- There is a need for increasing patient engagement during informed consent process
- Multiple perspectives on important information for the informed consent process

Abstract

One of the significant modifications to the Common Rule is the requirement that prospective participants are given information sufficient for a “reasonable person”. However, there is limited research on what types of information patients, caregivers, and providers consider “key information”. Although certain aspects of informed consent (IC) may be considered standard, considering individualized needs and preferences of patients is necessary for patient-centered

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