is a documented urgent medical need for the HCT/P, as defined in §1271.3(u).

(2) If you make an HCT/P available for use under the provisions of paragraph (d)(1) of this section, you must prominently label it "NOT EVALU-ATED FOR INFECTIOUS SUB-STANCES," and " WARNING: Advise patient of communicable disease risks." The following information must accompany the HCT/P:

(i) The results of any donor screening required under §1271.75 that has been completed;

(ii) The results of any testing required under §1271.80 or 1271.85 that has been completed; and

(iii) A list of any screening or testing required under §1271.75, 1271.80 or 1271.85 that has not yet been completed.

(3) If you are the establishment that manufactured an HCT/P used under the provisions of paragraph (d)(1) of this section, you must document that you notified the physician using the HCT/P that the testing and screening were not complete.

(4) In the case of an HCT/P used for an urgent medical need under the provisions of paragraph (d)(1) of this section, you must complete the donor-eligibility determination during or after the use of the HCT/P, and you must inform the physician of the results of the determination.

§1271.65 How do I store an HCT/P from a donor determined to be ineligible, and what uses of the HCT/ P are not prohibited?

(a) Storage. If you are the establishment that stores the HCT/P, you must store or identify HCT/Ps from donors who have been determined to be ineligible in a physically separate area clearly identified for such use, or follow other procedures, such as automated designation, that are adequate to prevent improper release until destruction or other disposition of the HCT/P in accordance with paragraph (b) or (c) of this section.

(b) Limited uses of HCT/P from ineligible donor. (1) An HCT/P from a donor who has been determined to be ineligible, based on the results of required testing and/or screening, is not prohibited by subpart C of this part from use 21 CFR Ch. I (4-1-06 Edition)

for implantation, transplantation, infusion, or transfer under the following circumstances:

(i) The HCT/P is for allogeneic use in a first-degree or second-degree blood relative;

(ii) The HCT/P consists of reproductive cells or tissue from a directed reproductive donor, as defined in §1271.3(1); or

(iii) There is a documented urgent medical need as defined in §1271.3(u).

(2) You must prominently label an HCT/P made available for use under the provisions of paragraph (b)(1) of this section with the Biohazard legend shown in §1271.3(h) with the statement "WARNING: Advise patient of communicable disease risks," and, in the case of reactive test results, "WARNING: Reactive test results, "WARNING: Reactive test results for (name of disease agent or disease)." The HCT/P must be accompanied by the records required under §1271.55.

(3) If you are the establishment that manufactured an HCT/P used under the provisions of paragraph (b)(1) of this section, you must document that you notified the physician using the HCT/P of the results of testing and screening.

(c) Nonclinical use. You may make available for nonclinical purposes an HCT/P from a donor who has been determined to be ineligible, based on the results of required testing and/or screening, provided that it is labeled:

(1) "For Nonclinical Use Only" and

(2) With the Biohazard legend shown in §1271.3(h).

§1271.75 How do I screen a donor?

(a) All donors. Except as provided under §1271.90, if you are the establishment that performs donor screening, you must screen a donor of cells or tissue by reviewing the donor's relevant medical records for:

(1) Risk factors for, and clinical evidence of, relevant communicable disease agents and diseases, including:

(i) Human immunodeficiency virus;

(ii) Hepatitis B virus;

(iii) Hepatitis C virus;

(iv) Human transmissible spongiform encephalopathy, including Creutzfeldt-Jakob disease;

(v) Treponema pallidum; and

(2) Communicable disease risks associated with xenotransplantation.