## Payment to Research Subjects -Information Sheet

## **Guidance for Institutional Review Boards and Clinical Investigators**

The Institutional Review Board (IRB) should determine that the risks to subjects are reasonable in relation to anticipated benefits [21 CFR 56.111(a)(2)] and that the consent document contains an adequate description of the study procedures [21 CFR 50.25(a)(1)] as well as the risks [21 CFR 50.25(a)(2)] and benefits [21 CFR 50.25(a)(3)]. It is not uncommon for subjects to be paid for their participation in research, especially in the early phases of investigational drug, biologic or device development. Payment to research subjects for participation in studies is not considered a benefit, it is a recruitment incentive. Financial incentives are often used when health benefits to subjects are remote or non-existent. The amount and schedule of all payments should be presented to the IRB at the time of initial review. The IRB should review both the amount of payment and the proposed method and timing of disbursement to assure that neither are coercive or present undue influence [21 CFR 50.20].

Any credit for payment should accrue as the study progresses and not be contingent upon the subject completing the entire study. Unless it creates undue inconvenience or a coercive practice, payment to subjects who withdraw from the study may be made at the time they would have completed the study (or completed a phase of the study) had they not withdrawn. For example, in a study lasting only a few days, an IRB may find it permissible to allow a single payment date at the end of the study, even to subjects who had withdrawn before that date.

While the entire payment should not be contingent upon completion of the entire study, payment of a small proportion as an incentive for completion of the study is acceptable to FDA, providing that such incentive is not coercive. The IRB should determine that the amount paid as a bonus for completion is reasonable and not so large as to unduly induce subjects to stay in the study when they would otherwise have withdrawn. All information concerning payment, including the amount and schedule of payment(s), should be set forth in the informed consent document.

Also see these FDA Information Sheets: "A Guide to Informed Consent Documents" (/RegulatoryInformation/Guidances/ucm126431.htm) "Recruiting Study Subjects." (/RegulatoryInformation/Guidances/ucm126428.htm) More in <u>Search for FDA Guidance Documents</u> (/RegulatoryInformation/Guidances/default.htm)

FDA Guidance Documents: General and Cross-Cutting Topics (/RegulatoryInformation/Guidances/ucm122044.htm)

Advisory Committee Guidance Documents (/RegulatoryInformation/Guidances/ucm122045.htm)

Clinical Trials Guidance Documents (/RegulatoryInformation/Guidances/ucm122046.htm)

<u>Combination Products Guidance Documents</u> (/RegulatoryInformation/Guidances/ucm122047.htm)

Import and Export Guidance Documents (/RegulatoryInformation/Guidances/ucm122048.htm)

International Council for Harmonisation (ICH) Guidance Documents (/RegulatoryInformation/Guidances/ucm122049.htm)

<u>Veterinary International Conference on Harmonization (VICH) Guidance Documents</u> (/RegulatoryInformation/Guidances/ucm122050.htm)