FAST FACTS

ALLIANCE A212102 - BLINDED REFERENCE SET FOR MULTICANCER EARLY DETECTION BLOOD TESTS

Eligibility criteria

Eligibility Criteria for Participants with a Cancer Diagnosis
1. Documentation of Disease:
   a. Histologic Documentation: Histologically confirmed diagnosis of invasive cancer
      • Stage: Stage I-IV per AJCC 7th edition, with the exception of patients with leukemia, lymphoma, and multiple myeloma.
         • For leukemia: Type (CLL, CML, ALL, AML)
         • For lymphoma: Stage I-IV based on Ann Arbor staging
         • For multiple myeloma: Stage I, II, III based on Revised International Staging System (RISS)
   b. One of the following tumor types:
      • Colorectal
      • Bladder
      • Head and Neck
      • Hepatobiliary
      • Lung
      • Lymphoma
      • Leukemia
      • Ovary*
      • Pancreas*
      • Multiple Myeloma
      • Gastric, esophageal or gastroesophageal
      • Breast
      • Thyroid
      • Kidney*
      • Endometrium
      • Prostate
      • Melanoma*
      • Sarcoma
      *For these specific cancer types only, patients may be enrolled prior to histologic confirmation of malignancy. Sites are required to contact the Study Chairs to review appropriateness for enrollment. See Section 4.3.2. Eligibility criteria for these patients are listed in Section 3.4.
2. No prior definitive systemic or local anti-cancer intervention.
3. Age ≥ 40 and ≤ 75
4. No known current pregnancy by self-report
5. No known or prior history of in situ or invasive malignancy (excluding in situ non-melanoma skin cancers) other than the current cancer diagnosis.
6. Willingness to provide blood samples for research use.
7. Absence of medical contraindications to a research blood draw volume of 60mL
8. No history of organ transplantation
9. Ability to read and comprehend English or Spanish

Eligibility is restricted to individuals who can comprehend and read English or Spanish given that participation in the study will require the ability to read and complete questionnaires that are available only in those two languages.

Eligibility Criteria for Participants without a Cancer Diagnosis and without Suspicion of Cancer

1. Age ≥ 40 and ≤ 75
2. No known current pregnancy by self-report
3. No known or prior history of in situ or invasive malignancy (excluding in situ non-melanoma skin cancers)
4. Willingness to provide blood samples for research use
5. Absence of medical contraindications to a research blood draw volume of 60mL
6. No history of organ transplantation
7. Ability to read and comprehend English or Spanish

Eligibility Criteria for Participants with a High Suspicion of Cancer

1. High suspicion of ovarian cancer, pancreatic cancer, kidney cancer, or melanoma by clinical and/or radiological assessment, with plans for histologic or cytologic confirmation within 28 days after study blood draw.
   • Examples of highly suspicious cases include: elevated CA125 and abnormal transvaginal ultrasound, suspicious renal or pancreatic mass on imaging, suspicious cutaneous lesion concerning for melanoma.
2. Central review of radiology reports and/or clinical documentation conducted by Study Chairs (see Section 4.3.2).
3. Age ≥ 40 and ≤ 75
4. No known current pregnancy by self-report
5. No known or prior history of in situ or invasive malignancy (excluding in situ non-melanoma skin cancers)
6. Willingness to provide blood samples for research use
7. Absence of medical contraindications to a research blood draw volume of 60mL
8. No history of organ transplantation
9. Ability to read and comprehend English or Spanish

Eligibility is restricted to individuals who can comprehend and read English or Spanish given that participation in the study will require the ability to read and complete questionnaires that are available only in those two languages.
**Schema**

At the time of registration**

- Patient consented and deemed eligible per applicable criteria
- Collection of biospecimens and data
- Registration (Step 1)**
- Collection of data

12 months (+/- 60 days) after registration

* Slot reservation required (see Section 4.3).

** For patients with a cancer diagnosis, biospecimens and data should be collected prior to any definitive therapy for the cancer.