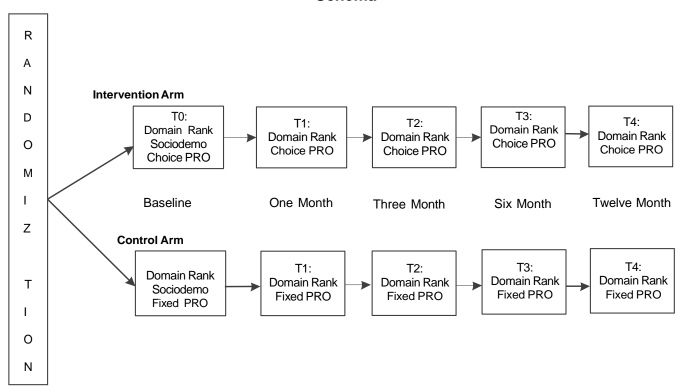


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Schema



Eligibility:	Randomization:	Domain Rank:
-Age 18 to 39	Stratified by sex,	Participant Ranks Domain by personal priority at each time point
-Within 12 weeks of	race, ethnicity, and	Fixed PRO:
diagnosis	age (emerging adults	PROMIS Global, PROMIS standard AYA 5 domains, Common
-Performance Status 0-3	18-25-year-old vs	Items
-Any stage of cancer	young adults 26-39-	Choice PRO:
-Favorable prognosis	year-old)	PROMIS Global, 5 ranked AYA domains, Common Items

Accrual Goal = 400

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3. Selection

3.1 Selection of Sites

All interested NCTN, LAPS, and NCORP sites will be invited to participate. There will be no specific exclusion criteria (i.e., minimum number of cases) because we would like to include a wide variety of different systems to better understand the variability in site-level AYA-specific resources and AYA patient volume. To ensure patient diversity in this study, all Minority/Underserved National Cancer Institute Community Oncology Research Programs (MU NCORP) sites will be specifically invited to participate.

3.2 Selection of Patients

Each of the criteria in the checklist that follows must be met in order for a patient to be considered eligible for this study. Use the checklist to confirm a patient's eligibility. For each patient, this checklist must be photocopied, completed and maintained in the patient's chart.

In calculating days of tests and measurements, the day a test or measurement is done is considered Day 0. Therefore, if a test is done on a Monday, the Monday four weeks later would be considered Day 28.

ECOG-AC	RIN Patient No.				
Patient's Initials (L, F, M) _					
Physician Signature and Date					
NOTE:	CTEP Policy does not allow for the issuance of waivers to any protocol specified criteria (http://ctep.cancer.gov/protocolDevelopment/policies_deviations.htm). Therefore, all eligibility criteria listed in Section 3 must be met, without exception. The registration of individuals who do not meet all criteria listed in Section 3 can result in the participant being censored from the analysis of the study, and the citation of a major protocol violation during an audit, and require reporting to the IRB of record as non-compliance.				
	Il questions regarding clarification of eligibility criteria must be directed to the roup's Executive Officer (<u>EA.ExecOfficer@jimmy.harvard.edu</u>) or the roup's Regulatory Officer (<u>EA.RegOfficer@jimmy.harvard.edu</u>).				
NOTE:	Institutions may use the eligibility checklist as source documentation if it has been reviewed, signed, and dated prior to registration/randomization by the treating physician.				
3.3 Eligibility Criteria					
3.3.	Patient must be ≥ 18 years and ≤ 39 years of age at registration.				
3.3.2	Patient must have a histologically confirmed diagnosis of primary cancer of any stage within 12 weeks (84 days) at registration.				
3.3	.3 Patient must not have a recurrence or second primary cancer.				
3.3	.4 Patients must not have basal cell skin carcinoma.				

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;	3.3.5	Patient must have received, be currently receive treatment for cancer, including su and/or radiation therapy.		
;	3.3.6	Patient must have an ECOG performance	e status 0-3.	
;	3.3.7	Patient must have a life expectancy >24 months.		
;	3.3.8	Patient must be able to complete questionnaires in English.		
;	3.3.9	Patient must have internet access throug smartphone.	h computer, tablet, or	
 ;	3.3.10	Patient must have an email address.		
;	3.3.11	Patient must have a mobile phone able with text messaging capabilities.		
;	3.3.12	Patient must be able to accurately provide self-report data (e.g. per clinical judgment, cognitive function is intact).		
;	3.3.13	Patient must be able to provide informed	consent.	
		Physician Signature	Date	

OPTIONAL:

This signature line is provided for use by institutions wishing to use the eligibility checklist as source documentation.