# Section: Contact Information

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# Section: Project Requirements and Description

#### Group: Requirements to submit AOI

A comprehensive review of previously published data has been completed. : Yes

The specific aims are clear and focused. : Yes

The investigator has appropriate experience and expertise to develop the concept proposal; if not, has identified a mentor or senior co-investigator. : **Yes** 

The investigator agrees to develop an initial draft of the concept proposal within 6 weeks of approval of the AOI and to finalize the concept proposal within 6 months. : **Yes** 

Project Title : Socioeconomic and Rural/Urban Differences in Adverse Outcomes Among Childhood Cancer Survivors

Planned research population (eligibility criteria):

The study population will include survivors in the expansion CCSS cohort who had complete information for socioeconomic status and zip code in the baseline questionnaire.

Proposed specific aims:

Aim #1: To evaluate differences in overall and cause-specific mortality by household income

Aim #2: Evaluate differences in overall and cause-specific mortality by rurality

# Aim #3: Assess effect modification of the relationship between mortality and rurality by household income

Will the project require non-CCSS funding to complete? : No

If yes, what would be the anticipated source(s) and timeline(s) for securing funding?:

#### Group: Does this project require contact of CCSS study subjects for:

Additional self-reported information: No

Biological samples : **No** Medical record data : **No** 

If yes to any of the above, please briefly describe. :

Group: What CCSS Working Group(s) would likely be involved? (Check all that apply)

Second Malignancy: Chronic Disease:

Psychology / Neuropsychology:

Genetics:

Cancer Control:

Epidemiology / Biostatistics : Primary

### Section: Outcomes or Correlative Factors

Late mortality: Primary

Second Malignancy: Secondary

#### Group: Health Behaviors

Tobacco: Alcohol:

Physical activity:
Medical screening:

Other:

If other, please specify:

# Group: Psychosocial

Insurance: Correlative Factors

Marriage:

Education : Correlative Factors
Employment : Correlative Factors

Other:

If other, please specify:

#### **Group: Medical Conditions**

Hearing/Vision/Speech:

Hormonal systems:

Heart and vascular: Secondary

Respiratory:
Digestive:

Surgical procedures:

Brain and nervous system:

Other:

If other, please specify:

# **Group: Medications**

Describe medications:

# Group: Psychologic/Quality of Life

BSI-18 : SF-36 :

CCSS-NCQ:

PTS : PTG : Other : If other, please specify:

#### Group: Other

Pregnancy and offspring:

Family history:

Chronic conditions (CTCAE v3): Secondary

Health status : Secondary

Group: Demographic

Age : Correlative Factors

Race : Correlative Factors

Sex : Other :

If other, please specify:

#### Group: Cancer treatment

Chemotherapy: Radiation therapy:

Surgery:

#### Section: Anticipated Sources of Statistical Support

CCSS Statistical Center: Yes Local institutional statistician:

If local, please provide the name(s) and contact information of the statistician(s) to be involved. :

Will this project utilize CCSS biologic samples? : No

If yes, which of the following?:

If other, please explain:

# Section: Other General Comments

Other General Comments:

This proposed analysis would be mentored/supervised by Dr. Smita Bhatia.

I agree to share this information with St. Jude: Yes