





### Feasibility of collecting survey-based patient reported outcome measures (PROMs) from patients living with advanced cancer: Emerging findings from the Living with Colorectal Cancer study

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**Interim Results** 

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**Presenter Disclosure** 

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Potential for conflict(s) of interest: None



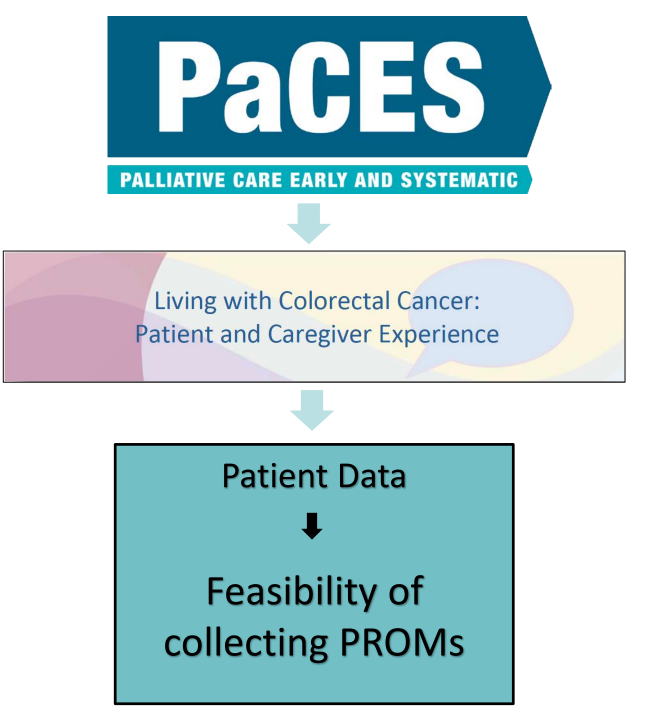




**Mitigating Potential Bias** 

Not Applicable





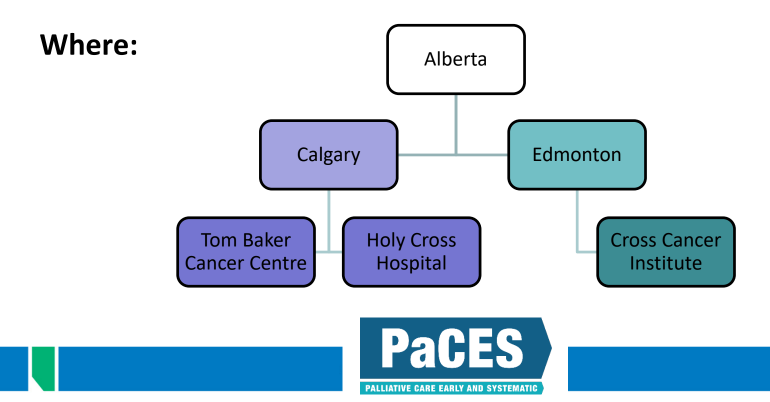
#### **Objective**:

- Measure how quality of life in this population changes over time
- Before and after the Palliative Care Early and Systematic (PaCES) pathway is implemented

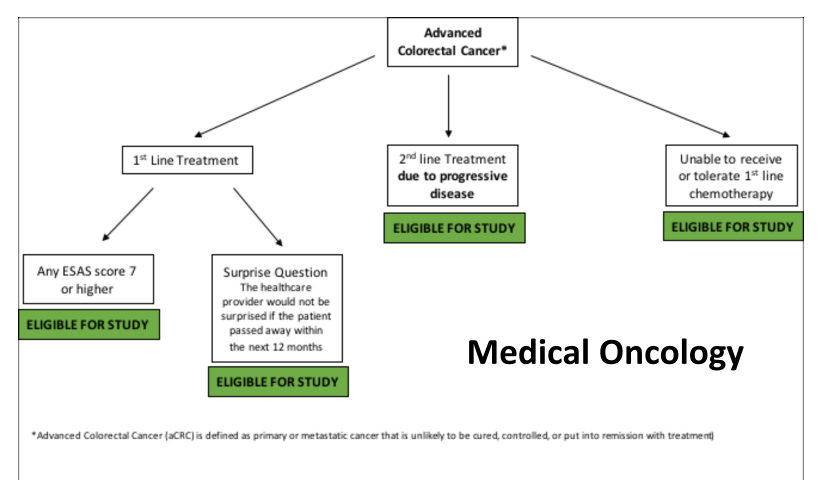


#### What:

- Observational study
- Gather experience of **patients** with advanced colorectal cancer and their caregivers



#### Who:





#### **Duration**:





#### **Types of participation**

### Full

- Patients complete surveys monthly

#### Partial

- Patients complete one set of surveys at enrollment only
- Following enrollment, study staff has access to the patient's medical record but patients are not required to complete any further surveys



#### Methods:

- Initial and periodic PROMs and PREMs
  - <u>PROM</u>: capture a patient's perception of **their health**
  - <u>PREM</u>: capture a patient's perception of their experience with health care or a service



	Please check the box that best describes how you feel <b>NOW.</b>		
Τος	I feel at peace.		
	<ul> <li>1. Not at all</li> <li>2. A little bit</li> <li>3. A moderate amount</li> </ul>		
PF	<ul> <li>3. A moderate amount</li> <li>4. Quite a bit</li> <li>5. Completely</li> </ul>		

What concerns have you had in the last month? Check any boxes that have concerned you.								
Emotional Fears/Worries Sadness Frustration/Anger Changes in appearance Intimacy/Sexuality Thoughts of ending my life	Physical         Fever/Chills         Bleeding/Bruising         Cough         Headaches         Concentration/Memory         Vision or hearing changes         Numbness/Tingling         Sensitivity to cold         Changes to skin/nails         Bladder problems         Lymphedema/Swelling         Range of motion         Strength         Speech difficulties	Nutrition         Weight gain (amount)         Weight loss (amount)         Special diet         Difficulty swallowing         Mouth sores         Taste changes         Heartburn/Indigestion         Vomiting         Diarrhea         Constipation         Informational         Understanding my illness and/or treatment         Talking with my health care team         Making treatment decisions         Knowing about available resources         Taking medications as prescribed						
Social/Family/Spiritual  Feeling alone Feeling like a burden to others Vorry about friends/family Support with children/partner Meaning/Purpose of life Faith								
Practical UVork/School Finances Getting to and from appointments Home Care	□ Sleep Mobility □ Dizziness □ Walking/Mobility □ Trouble with daily activities (e.g. bathing, dressing)							
<ul> <li>Accommodation</li> <li>Quitting tobacco</li> <li>Drug costs</li> <li>Health insurance</li> <li>How much alcohol you drink</li> </ul>	Other Concerns:							

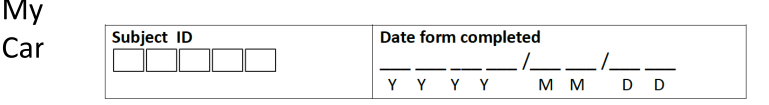
#### **CUMMING SCHOOL OF MEDICINE**



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#### MY CONVERSATIONS

#### Living with Colorectal Cancer: Patient and Caregiver Experience



Some people talk with their healthcare providers (e.g. doctors, nurses, etc.) about their illness, healthcare preferences, and planning for the future.

1. In the last month, has a healthcare provider talked with you about the following? (Check ( $\sqrt{}$ ) all that apply)

Asked you what is important to you as you consider your healthcare preferences (such as your values, wishes, goals, or spiritual beliefs)

Talked to you about your prognosis (life expectancy, predicted course or outlook of your illness)

Given you the opportunity to express your fears or to discuss what concerns you

Asked you about treatments you prefer to have or not have if you were to be very sick or at the end of life

□ I had contact with a healthcare provider in the last month but I had none of these conversations (Go to question 2)

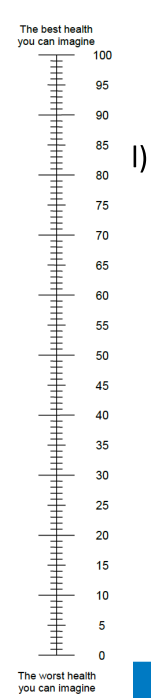
□ I did not have contact with a healthcare provider in the last month (Go to question 3)

#### **Tools used**

#### EQ-5D-5L-

- We would like to know how good or bad your health is TODAY.
- This scale is numbered from 0 to 100.
- 100 means the <u>best</u> health you can imagine.
   0 means the <u>worst</u> health you can imagine.
- Mark an X on the scale to indicate how your health is TODAY.
- Now, please write the number you marked on the scale in the box below.

YOUR HEALTH TODAY =

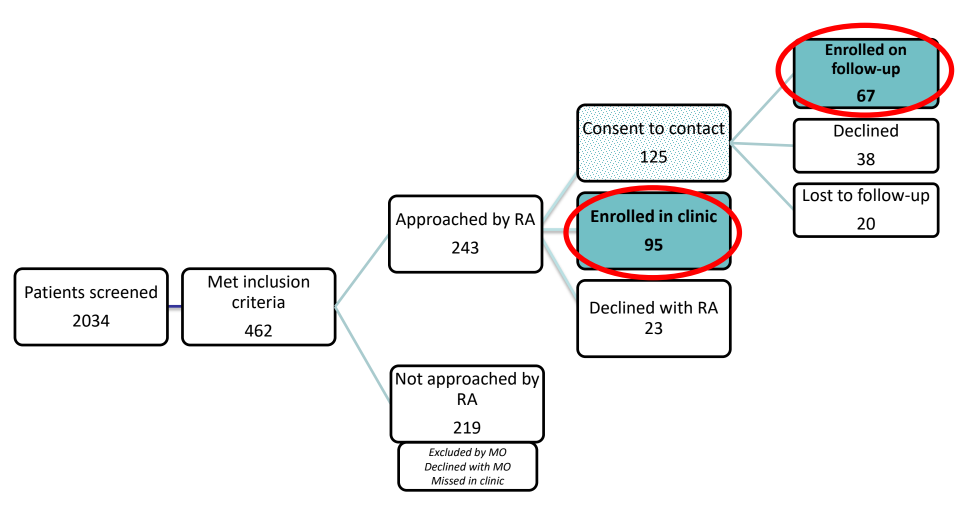


#### **Methods of participation**



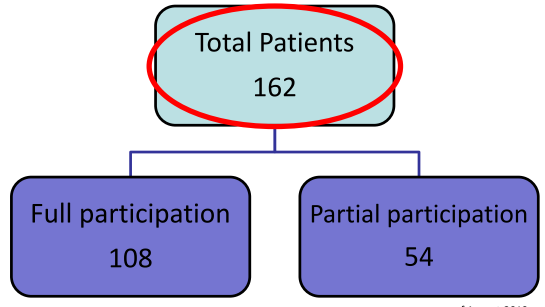


#### **Screening & Recruitment**





#### **Screening & Recruitment**



as of August 2019



#### **Demographics\***

Full Participants	Median age (Range)	Female	Male	Reported speaking a language other than English
108	64 (32-88)	46 (43%)	62 (57%)	21%

\* Full Participants only





#### **Non-English speaking patients**

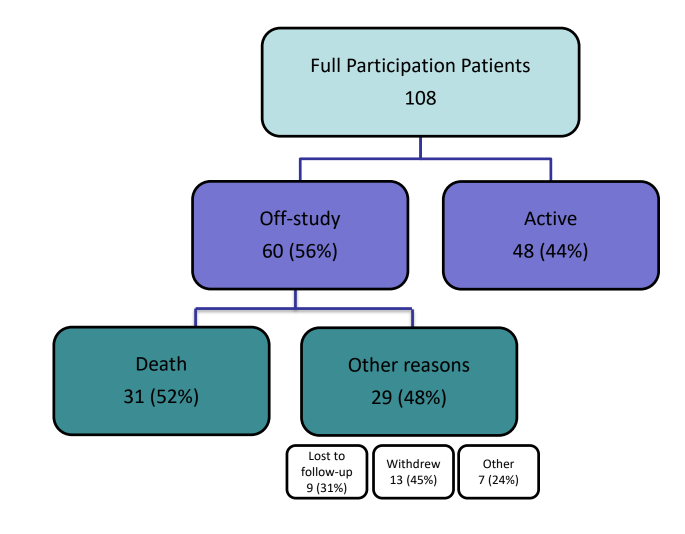
#### 12 eligible patients identified\*

- 11 declined
  - 3 too sick/ in hospice on follow-up
  - 1 deceased on follow-up
  - 4 no longer interested in participating
  - 2 not interested in participating because surveys must be done in person/via mail (no online option)
  - 1 off all treatment and moved to another country
- 1 enrolled (Cantonese speaking, Edmonton)

\*Arabic, Tagalog (x4), Spanish, Cantonese (x2), Vietnamese (x2), Russian (x2)



#### **Off-study rates**





#### **Off-study statistics\***

Days between enrollment date and off-study date:

- 40-569 days (range)
- 186 days (median)



\*Full participation patients only



#### **Missing Data**

Reasons for missing data (surveys that are not completed):

- Too sick or in hospital
- o Death
- o Lost to follow-up
- Patient requests to stop follow-up surveys
- o Vacation
- o Other
  - o -patient forgot, technical difficulties etc.

# $\frac{120}{906}$ = 15%



So what <u>is</u> the feasibility of collecting survey-based patient reported outcome measures (PROMs) from patients living with advanced cancer?





It is challenging but possible to engage and retain patients with advanced cancer in research focused on PROs.



Offering flexible modes of participation assists with patient engagement.



#### **Key Learnings**

### SUCCESSES

#### 1. Patient engagement

-Face-to-face contact appreciated -Patients eager to provide experiences

# 2. Multiple methods for survey completion

-Allows patients to participate in a manner most practical for their lifestyle

-Did not impact PROMs collection

#### CHALLENGES

# 1. Burden on healthcare providers

-Clinic flow: room occupied while Research Assistant recruits and enrolls patients

# 2. Maintaining long-term contact with patients

-Patients withdraw when they become too sick to participate





Determine the time between patients going off-study and actual date of death



#### **Future Considerations**

"It is interesting to think about my responses to the survey questions every month but it is pretty structured. I would love to tell my story from the beginning, starting with when and how I was diagnosed so you can get the full picture of my experience."

-Patient A

Some patients expressed a desire to provide more details about their experience

- Consider a qualitative survey/ data collection method
  - Dr. Maria Santana's companion study



#### **Future Considerations**

#### Explore other ways to engage non-English speaking patients

 Currently, patients must complete surveys via mail only (no online versions of the surveys in other languages)



"I'm happy to fill out these surveys every month and let you know how things are going with me. I hope this will help someone else in the future."

-Patient B

*"It's nice that I can do the surveys during my chemo treatment but can also do them with you over the phone if I'm not coming to the Cancer Centre that month."* 





### **Thank You**

### **Questions / Comments?**









Canadian Institutes Instituts de recherche of Health Research en santé du Canada