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# Implementing an Inclusive, Multidisciplinary Supportive Care Model to Provide Integrated Care to Breast and Gynaecological Cancer Survivors: A Case Study at an Asian Ambulatory Cancer Centre

INTEGRATED CARE  
CASE

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## ABSTRACT

**Introduction:** Supportive care models considering inclusivity and community services to improve integrated care for cancer survivors are limited. In this case study, we described the implementation of a multidisciplinary care model employing routine distress screening and embedded integrated care pathways to integrate care across disciplines and care sectors, while remaining inclusive of the multi-ethnic and multilingual population in Singapore. We reported implementation outcomes after 18 months of implementation.

**Description:** We reviewed the model's process indicators from September 2019 to February 2021 at the largest public ambulatory cancer centre. Outcomes assessed included penetration, fidelity to screening protocol, and feasibility in three aspects – inclusiveness of different ethnic and language groups, responsiveness to survivors reporting high distress, and types of community service referrals.

**Discussion/conclusion:** We elucidated opportunities to promote access to community services and inclusivity. Integration of community services from tertiary settings should be systematic through mutually beneficial educational and outreach initiatives, complemented by their inclusion in integrated care pathways to encourage systematic referrals and care coordination. A hybrid approach to service delivery is crucial in ensuring inclusivity while providing flexibility towards external changes such as the COVID-19 pandemic. Future work should explore using telehealth to bolster inclusiveness and advance community care integration.

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care; inclusive; COVID-19

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## INTRODUCTION

According to the National Cancer Institute, a person with a cancer diagnosis is termed a survivor from diagnosis to the end of life [1]. Supportive care models aim to promote integrated care provision to survivors through the delivery of supportive care services, such as psycho-oncology services, rehabilitation, and palliative care, to prevent and manage adverse effects from cancer and its treatment [2, 3]. Among cancer survivors, an integrative approach combining different health disciplines is necessary to effectively manage the myriad of physical, emotional, social, psychological, practical, and spiritual issues throughout survivorship [4, 5].

Typically, cancer survivors are managed under the oncologist-led model where oncologists serve as the predominant cancer-related care providers in the tertiary setting [6]. Hence, oncologists play the pivotal role in providing supportive care and initiating referrals to supportive care services [3]. Two major gaps exist in this oncologist-centric model that warrants a paradigm shift. First, there is a disproportionate focus on managing physical problems than psychosocial issues due to clinical time constraints, inadequate education and training, a lack of awareness of available services, and a lack of knowledge about how to integrate attention to psychosocial health needs into their practices [7–10]. Second, integrated care provision across disciplines and sectors is compromised with inadequate infrastructural support and a lack of clearly defined referral pathways available for oncologists [8, 11]. Consequently, cancer survivors' access to supportive care services is heterogenous, fragmented, and subjected to significant inter-oncologist variability [7]. Care fragmentation is further associated with reduced resource allocation efficiency, poorer symptom control, higher costs, and failure to deliver consistent and coordinated patient-centred care [12, 13].

Supportive care models employing routine distress or care needs screening with follow-up care are then increasingly advocated and implemented to allow a holistic assessment of cancer survivors' needs for survivor-centric care personalization [14–17]. The basis for the theoretical benefits of such models includes early identification, evaluation, and timely management of active problems. Recognizing the diversity of care needs across the survivorship continuum, these models tailor evidence-based interventions and coordinate multidisciplinary supportive services.

However, several knowledge gaps remain to be addressed. First, existing evidence on the effectiveness of such supportive care models are inconclusive due to poor reporting of implementation outcomes [18]. Second, integrated care provision needs to stay inclusive of survivors across different cultural backgrounds, languages, and health literacy levels to avoid potential

health disparities [19, 20]. However, there is limited literature describing inclusivity considerations during care design and inclusiveness is rarely evaluated as an outcome measure. Lastly, while there are increasing calls for community engagement to provide integrated cancer survivorship care across health sectors, inadequate cancer-related knowledge among community care providers and suboptimal care coordination across settings continue to pose as challenges [21]. In summary, there is a demand for robust evaluation of supportive care models implemented with considerations of inclusivity and access to community services to improve integrated care for cancer survivors.

Singapore is a high-resource country in Southeast Asia with a multi-ethnic and multilingual population. The design and launch of the Accessible Cancer Care to Enable Support for Cancer Survivors (ACCESS) supportive care model in National Cancer Centre Singapore (NCCS) was intended to address the abovementioned knowledge gaps. The ACCESS model employed an inclusive design and routine distress screening to triage cancer survivors with differing care needs. Specifically, embedded integrated care pathways serve as facilitating mechanisms to integrate care referrals to different health disciplines in the tertiary setting and integrate survivorship care across tertiary and primary care sectors. This study aims to describe the ACCESS implementation process and to report implementation outcomes after an 18-month period.

## METHODS

### STUDY DESIGN

This study is part of a pragmatic type I effectiveness-implementation hybrid study designed to evaluate the impact of the ACCESS model in an ongoing cluster randomized controlled trial (NCT04014309) while collecting information on its implementation [22]. The literature recommended this hybrid approach to collect both implementation and effectiveness data to develop an evidence base that could promote widespread intervention adoption [23].

We focused on reporting the implementation outcomes from September 2019 to February 2021 here. This implementation study was approved by the SingHealth Centralised Institutional Review Board (CIRB 2020/2789) with the waiver of informed consent. Reporting of implementation metrics followed the Standards for Reporting Implementation Studies, and the model was described according to the minimum recommended components [24, 25].

### SETTING AND TARGET POPULATION

NCCS is the largest comprehensive ambulatory cancer centre serving 65% of adult cancer patients in Singapore's

public sector [26]. Funding from Temasek Foundation Cares supported the scale of model implementation in half of the medical oncologist clinics seeing breast and gynaecological cancer survivors, the most common incident cancers in women locally [27, 28]. Cancer survivors were eligible if they were ≥21 years, diagnosed with breast or gynaecological cancer, and had received outpatient care from participating medical oncologists minimally once during the review period.

**THEORETICAL BASIS OF THE ACCESS MODEL**

The ACCESS model adopted the supportive care framework outlined by Fitch et al. (Figure 1) that described a tiered approach with two underlying principles [4, 17]. First, basic health information and distress and care needs screening were offered to cancer survivors at all medical oncologist visits, regardless of language preference, and literacy levels. Second, access to supportive care resources was then determined based on the level of distress and care needs.

**ACCESS MODEL IMPLEMENTATION DESIGN AND STRATEGIES**

Implementation of the ACCESS intervention was guided by the Consolidated Framework for Implementation, which outlined determinants of successful implementation in the following domains: individual survivor characteristics, intervention characteristics, processes, and the organizational setting [29]. Complementarily, implementation strategies were developed to maximize

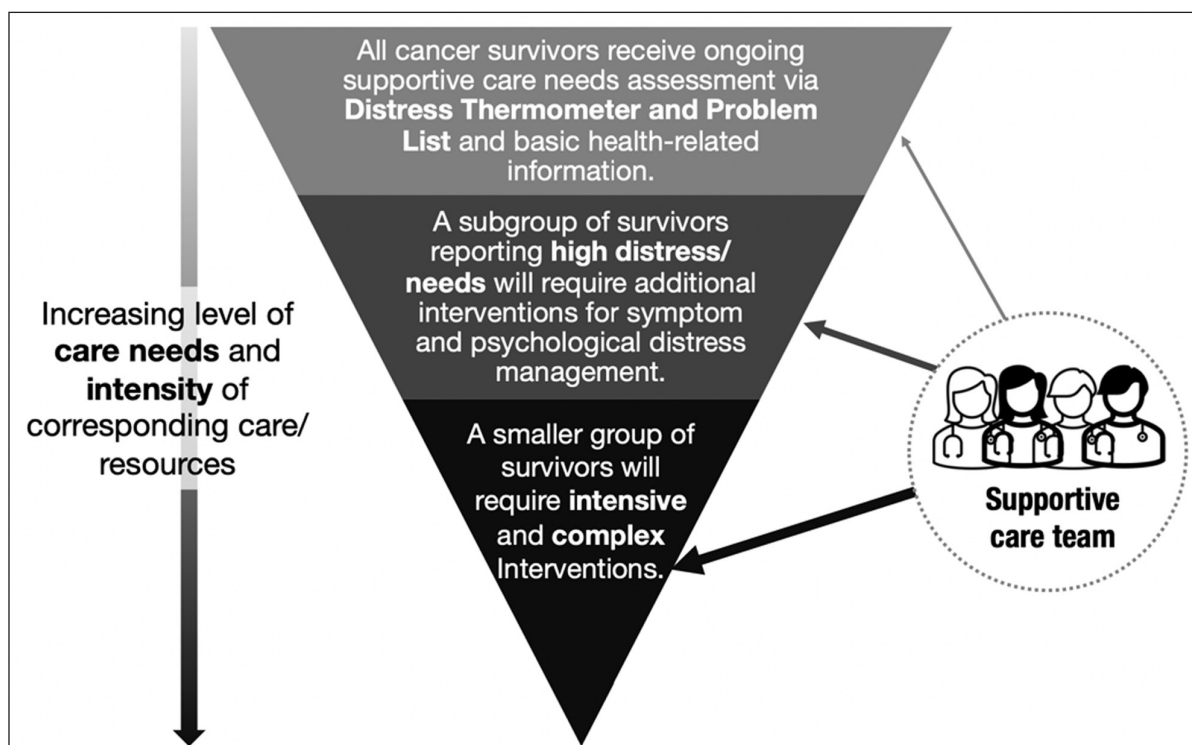
screening outreach, facilitate the timely identification of survivors with high care needs, and improve access to tertiary and community support services. Implementation strategies are methods or techniques used to enhance the adoption, implementation, and sustainability of a clinical program [30]. Table 1 details the employed implementation strategies according to reporting recommendations [30, 31].

**Core workgroup formation**

A multidisciplinary core workgroup that included oncologists, a rehabilitation physician, nurses, clinical pharmacists, and a psychologist oversaw the ACCESS model implementation. The workgroup met monthly to review implementation progress and anchored outreach efforts to relevant stakeholders (NCCS institution leaders, participating oncologists, funder, and the Singapore Ministry of Health), actively addressing implementation issues and elucidating analyses required for funding considerations to support long-term implementation.

**Education initiatives**

Collaborating with the NCCS Cancer Education and Information Services and corporate communications, the workgroup revamped, consolidated, and standardized available education and community resources on the institution website [32]. A leaflet summarizing the ACCESS model and containing a link/QR code to this institution website was provided to all survivors to empower them and their caregivers to self-manage



**Figure 1** Tiered care approach of the Accessible Cancer Care to Enable Support for Cancer Survivors model. Routine distress and care needs screening were provided to all cancer survivors before triaging to further follow-up by a supportive care team based on reported levels of care needs.

STRATEGY	CFIR DOMAIN	ACTORS AND ACTIONS	TARGET AUDIENCE	TEMPORALITY	JUSTIFICATION	ANTICIPATED IMPACT
Establish a core workgroup to ensure procedural accountability	Organizational (inner setting)	The core workgroup met regularly to discuss implementation challenges and elicit feedback for workflow streamlining.	Core workgroup, clinicians delivering the intervention, service coordinators	Monthly meetings were held with key process indicators reporting to the funder every quarter.	Facilitates iterative feedback and improvement process and brainstorms strategic engagement with stakeholders.	Enhanced feasibility and sustained intervention adoption.
Develop educational initiatives	Process	The core workgroup collaborated with the institutional education department to develop informational resources and plan educational activities.	Cancer survivors and caregivers, community service partners	Educational activities were concurrent with intervention implementation.	Promotes awareness of intervention and self-management by survivors and network weaving with community partners through educational outreach efforts.	Improved adherence to screening, increased number of trained community partners, and increased referrals to community services.
Strengthen the inclusivity of screening procedures	Individual survivor characteristics, intervention characteristics, process	The core workgroup culturally adapted the screening tool for the local population and translated it into multiple languages. Informational system and trained service coordinators supported a hybrid screening administration mode.	Cancer survivors	Translation occurred before the intervention launch. Efforts to improve the informational system for EMR integration are ongoing.	Targets survivor-level language and health literacy barriers to enhance screening uptake and adherence.	No systematic differences in screening completion rates across survivors of different ethnicities and health literacy levels.
Develop standardized, integrated care pathways	Intervention characteristics, process	The core workgroup led the development of standardized care pathways to map evidence-based interventions to all reported problems in the screening tool.	Supportive care team members	Pathways were drafted before intervention launch and amenable to changes following workgroup meetings.	Reduces variation in clinical practice across supportive care team members and provides systematic access to appropriate service referrals.	High responsiveness to survivors reporting high distress levels.

**Table 1** Summary of implementation strategies employed in the Accessible Cancer Care to Enable Support for Cancer Survivors intervention.

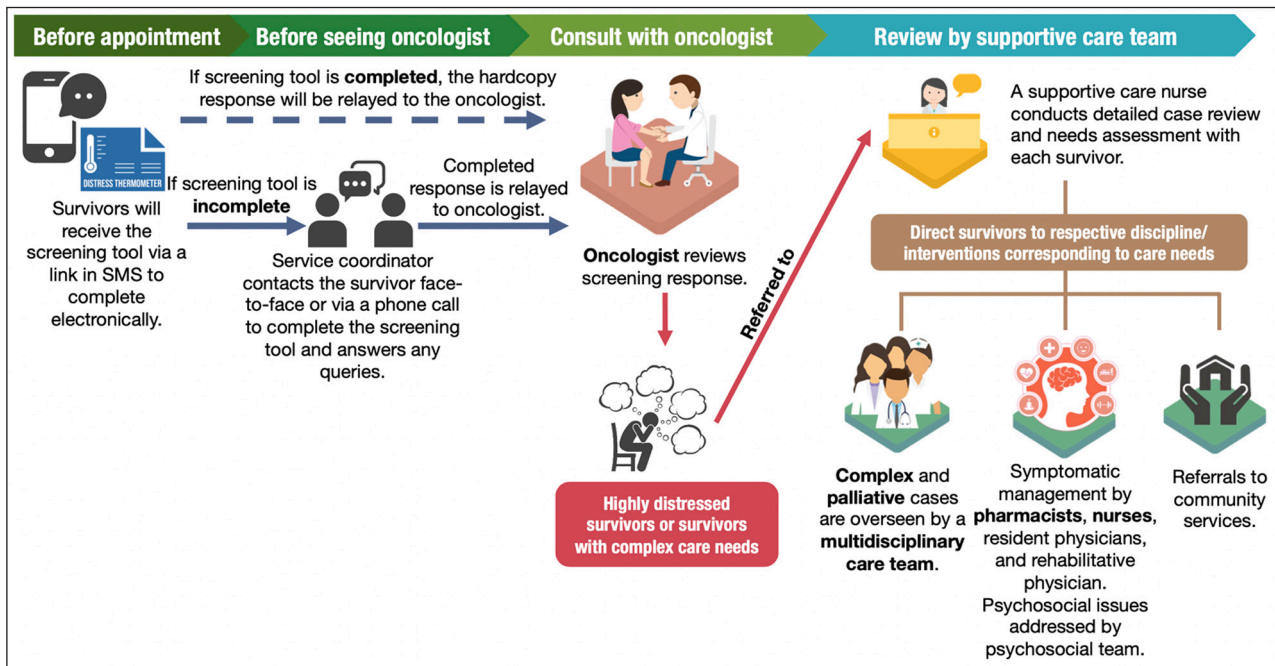
Abbreviations: CFIR, Consolidated Framework for Implementation Research; EMR, electronic health records.

problems, promoting self-efficacy [33]. Furthermore, to engage community partners in cancer survivorship care delivery, the workgroup gathered expertise from multiple disciplines to plan symposiums and lead educational activities, including experiential clinic attachments and curriculum development. Through these outreach efforts, an extensive list of available community services was consistently updated and mapped to survivorship care needs, facilitating downstream systematic referrals based on survivors' reported problems.


### Inclusive screening procedures

Trained service coordinators with science-related degrees facilitated the screening process (Figure 2). The ACCESS model was introduced to all survivors at their first visit during the review period. Subsequently, before

each medical oncologist visit, survivors received unique webpage links to the screening tool (Distress Thermometer and Problem List, DTPL) via short messaging services. The original English version was culturally adapted from the National Comprehensive Cancer Network (Figure 3) [17]. Survivors rated their distress levels on a Likert scale of 0-10 and indicated the problems (physical, practical, family, emotional, and spiritual) experienced in the past week. Self-administration typically took 5-10 minutes and responses were captured electronically. Several strategies were adopted to maximize inclusivity. First, the DTPL was translated into other official languages of Singapore (Mandarin, Malay, or Tamil) to accommodate the multilingual population. Second, on the day of clinic visit, service coordinators either approached or contacted survivors who did not complete the screening



**Figure 2** Overview of the clinical workflow for the screening procedures using the Distress Thermometer and Problem List, discussion during oncologists’ review, and subsequent management of survivors referred to the supportive care team.



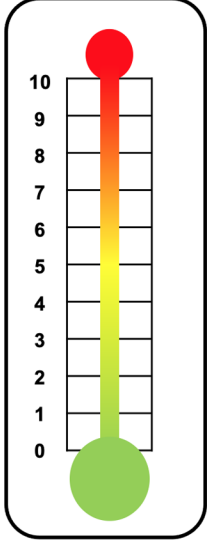
## NCCS

### Distress Screening

**DISTRESS THERMOMETER**

Please circle below the number (0-10) that best describes how much distress\* you have been experiencing in the past 1 week, including today:  
\*Distress= physical / mental strain and stress

You do not have to fill this up if you have done it within the last 1 month or if nothing has changed from your last input.



Version 1, 2018. Adapted from NCCN Guidelines V.2.2018 Distress Management.

**PROBLEM LIST**

Please tick if any of the following has been a problem or concern for you in the past 1 week, including today.

**Physical Problems**

- Appearance
- Breathing
- Problems in urination
- Constipation
- Diarrhoea
- Appetite
- Feeling tired
- Abnormal swelling in legs or arms
- Fevers
- Moving around
- Indigestion
- Memory / Concentration
- Mouth / Swallowing problems
- Nausea / Vomiting
- Nose dry / congested
- Pain
- Sexual problems
- Hair / Skin problems
- Sleep problems
- Tingling / numbness in feet / hands
- Hot flashes (feeling excessively hot / sweaty)
- Speech / Hearing problems

**Information needs**

- Specify: \_\_\_\_\_
- Other problems not listed; specify: \_\_\_\_\_

**Practical Problems**

- Caring responsibilities (child, parents, etc)
- Financial (include insurance)
- Transportation
- Work / School
- Treatment decision
- Household chores
- Bathing / Dressing / Toileting

**Family / relationship Problems**

- Relationship with family / friends
- Ability to have children
- Family health issues

**Emotional Problems**

- Sadness / Depression
- Worry / Fear / Anxiety
- Loss of interest in usual activities
- Anger / Frustration / Guilt
- Hopelessness
- Loneliness / Isolation

**Spiritual / Religious Concerns**

- Loss of faith or other spiritual concerns
- Loss of meaning / purpose of life
- Not at peace / feeling regret about past

**Figure 3** Distress Thermometer and Problem List adapted from the National Comprehensive Cancer Network.

tool electronically beforehand to administer the tool and address usability concerns. Lastly, hardcopy of the DTPL was also available for survivors without access to phones or internet.

**Clinical responses to screening results (Figure 2)**  
All screening responses were physically relayed to oncologists for review and discussion during consultations. Survivors were referred to the supportive

care team within the same day when (i) survivors reported a DT score  $\geq 6$ , indicative of high distress, and/ or (ii) additional review was deemed necessary based on oncologist’s discretion, regardless of DT score. A higher DT score of 7 was selected initially and revised to 6 after two months of implementation based on manpower capacity considerations. Both scores could identify participants with clinically significant distress in the local population [34]. All highly distressed survivors were offered a review with the supportive care team in-person or through telehealth.

**Review by a multidisciplinary supportive care team using integrated care pathways (Figure 4)**

Typically, a supportive care nurse would first conduct detailed needs assessments for each referred survivor before providing care according to standardized integrated care pathways. These care pathways were available for all problems reflected on the DTPL to ensure that screening results lead to actionable interventions. The workgroup developed these pathways by locally adapting interventions and recommendations from guidelines to reduce clinical variation when addressing reported supportive care needs. The nurse then facilitated access to other team members from diverse health disciplines (e.g., pharmacy, psychology, social work, rehabilitation) and community services. A further

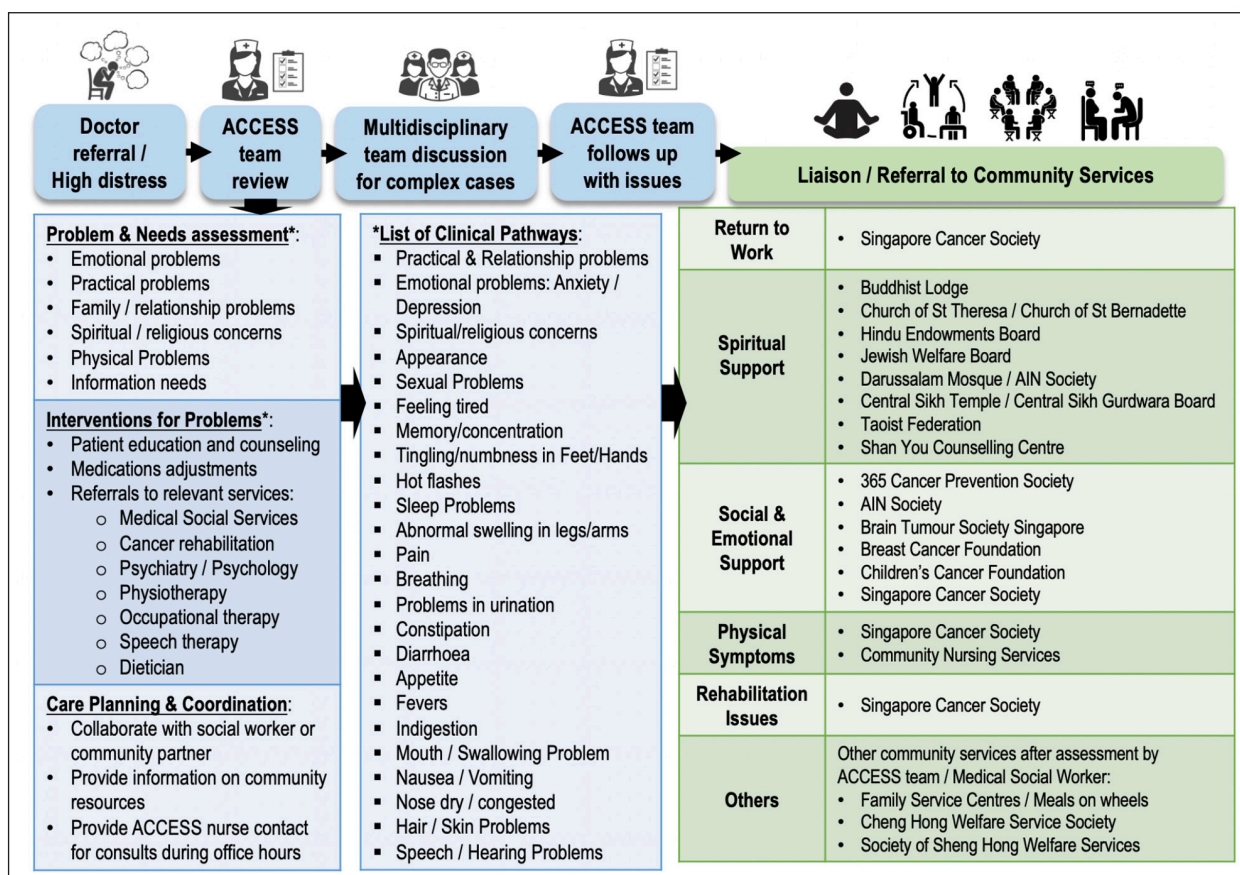
subgroup of survivors who presented with complex needs or palliative concerns were referred to multidisciplinary care team biweekly reviews.

**IMPACT OF COVID-19**

Responding to COVID-19, Singapore raised its disease outbreak alert to orange in February 2020, enforcing control measures and mobility restrictions [35]. A lockdown was imposed from 7 April to 1 June 2020 before gradual reopening [36]. Workflows were modified to adhere to institutional guidelines during periods of tightened restrictions. First, non-emergency visits were postponed. Second, all in-person encounters between service coordinators and survivors were replaced by calls. Third, relaying of screening results to oncologists was impermissible during the lockdown. In contrast, supportive care team workflows were minimally disrupted as telehealth consult was already available.

**OUTCOME EVALUATION**

We assessed the following implementation outcomes based on Procter et al.’s taxonomy [37]. We determined the penetration of our model by quantifying the number of survivors introduced to the ACCESS model and those who received self-help educational resources. Next, we assessed fidelity by tabulating adherence rates to the screening protocol based on the proportion of attended



**Figure 4** Overview of supportive care interventions and a network of community services referrals available for survivors based on care needs.

clinic visits with corresponding screening responses. Adherence rates were categorized into: <50%, reflecting adherence less than half of the time; 50–99%, reflecting adherence most of the time; and 100%, reflecting complete adherence.

The feasibility of implementing the model in real-world settings was assessed in three aspects. First, the inclusivity of the screening procedures was evaluated by determining whether screening tool completion rates differed by ethnicity and language preferences. The mode of screening tool completion was summarized as a proxy to measure the inclusiveness of survivors with lower literacy or who were less technology-savvy. Second, the proportion of screened survivors reporting high distress was determined. Responsiveness to these survivors was evaluated by tabulating the proportion of highly distressed survivors who received access to the supportive care team. Third, the access to community services was described by the number and types of referrals made to community services and corresponding acceptance and attendance rates. Lastly, implementation cost was assessed by the additional manpower required to support the screening procedures which were not part of usual care. Additional subgroup analyses were conducted to assess the impact of COVID-19 restrictions on penetration, mode of screening, and screening completion rates.

### SAMPLE SIZE CALCULATION AND STATISTICAL ANALYSIS

Based on past clinical load, we expected 3500 eligible survivors within the 18-month review period. All data were analysed using STATA 17. Descriptive statistics were used to describe survivors' characteristics and outcome indicators. Mean and standard deviation were used to summarize continuous variables, while counts and percentages were used for categorical variables. A multivariable Poisson regression model was used to analyse screening tool completion rates by ethnicity and language preferences. The model was further adjusted for survivors' age where adherence rates among the adolescent and young adult group were significantly lower than expected compared to older survivors in a study that evaluated distress screening outcomes across Commission on Cancer-accredited cancer programs [38]. All statistical tests were 2-sided with  $\alpha = .05$ .

## RESULTS

### PENETRATION

Among 2357 eligible breast and gynaecological cancer survivors, we successfully introduced the ACCESS model and screened 1853 (78.6%) survivors minimally once. Screened survivors had mean (SD) age of 60.7 (11.2) years, were predominantly Chinese (1496/1853,

80.7%) and were mostly diagnosed with breast cancer (1683/1853, 90.8%) (Table 2). Self-help educational resources were distributed to 1518 (81.9%) survivors.

### FIDELITY AND IMPLEMENTATION COST

Overall, the tasks and time required to support screening amounted to service coordinators operating at two full-time equivalents (Table 3). The overall adherence rate to the screening protocol was 77.4%, with 5779 screening responses obtained from 7464 visits. For 1259 survivors with multiple visits during the review period, 412 (32.7%) survivors completed the screening tool at between 50–99% of their visits, while 683 (54.3%) survivors demonstrated complete adherence (Table 4).

### FEASIBILITY – INCLUSIVITY

Most screening responses captured were completed by survivors electronically (67.6%) without assistance by service coordinators. The English version was most used while 22.1% preferred non-English version. Completion rates were not found to differ by language preferences, before and adjusting for age and ethnicity (Table 5). Among ethnic groups, the screening tool completion rate was 15% lower among Malays [adjusted rate ratio (95% CI): 0.85 (0.76, 0.95),  $P = 0.004$ ] than in Chinese, after adjusting for age and language preference.

### FEASIBILITY – RESPONSIVENESS

A total of 529/1853 (28.6%) screened survivors reported high distress minimally once during the review period. Among them, 231 (43.7%) survivors only reported high distress at repeat visits. Responding to this group, 19 (3.6%) survivors were uncontactable despite three

CHARACTERISTIC	N (%)
Age, mean (SD)	60.7 (11.2)
Ethnicity	
Chinese	1496 (80.7%)
Malay	186 (10.0%)
Indian	115 (6.2%)
Others	56 (3.0%)
Cancer diagnosis	
Breast	1683 (90.8%)
Gynaecological	170 (9.2%)
Language preference for screening tool	
English	1444 (77.9%)
Mandarin	385 (20.8%)
Malay	21 (1.1%)
Tamil	3 (0.2%)

**Table 2** Characteristics of ACCESS model recipients (N = 1853).



TASK	TIME REQUIRED	FREQUENCY	TOTAL TIME REQUIRED PER WEEK (MINUTE)
Export a list of scheduled visits in the upcoming week	30 minutes for all patients	Weekly	30
Register survivors in the electronic system before DTPL could be disseminated	5 minutes per survivor	Weekly	750
Schedule DTPL for the electronic system to trigger a SMS with the link for completion to survivors a few days before scheduled visits	10 minutes per survivor	Weekly	1500
Download DTPL responses from the electronic system into a compiled Excel spreadsheet	5 minutes per download	Daily	25
Transpose compiled responses into individual responses	5 minutes per response	Daily	750
Relay screening results to oncologists in clinics before consults physically and assist survivors with in-person completion of screening tool	Full clinic hours (6 hours daily)	Daily	1800
<b>Total</b>	4855 (equivalent to approximately <b>2 FTE</b> )		

**Table 3** A summary of tasks and time required to support screening procedures weekly based on an estimated caseload of 150 survivors per week.

Abbreviations: DTPL, Distress Thermometer and Problem List; FTE, full-time equivalent.

NUMBER OF VISITS IN THE REVIEW PERIOD	COMPLETION RATE		
	<50%, N (%)	50–99%, N (%)	100%, N (%)
All survivors with multiple visits	164 (13.0%)	412 (32.7%)	683 (54.3%)
3 visits (n = 171)	28 (16.4%)	46 (26.9%)	97 (56.7%)
4 visits (n = 132)	9 (6.8%)	55 (41.7%)	68 (51.5%)
5 visits (n = 119)	17 (14.3%)	35 (29.4%)	67 (56.3%)

**Table 4** Screening completion rates among survivors with multiple medical oncology visits during the 18-month implementation period (N = 1259).

VARIABLES	UNADJUSTED COMPLETION RATE RATIO (95% CI)	ADJUSTED COMPLETION RATE RATIO (95% CI)
Ethnicity		
Chinese	Reference	Reference
Malay	0.89 (0.79, 0.99)	0.85 (0.76, 0.95)
Indian	1.01 (0.91, 1.12)	0.99 (0.90, 1.10)
Others	1.00 (0.87, 1.14)	0.98 (0.85, 1.14)
Preferred language		
English	Reference	Reference
Non-English	0.96 (0.90, 1.00)	0.95 (0.89, 1.02)
Age <sup>a</sup>	1.00 (0.99, 1.00)	1.05 (0.99, 1.00)

**Table 5** Poisson regression analysis of screening completion rates for survivors with multiple medical oncology visits (N = 1259).

<sup>a</sup>Modelled as a continuous variable.

separate call attempts on different days while attempts to contact another 89 (16.8%) survivors are ongoing. Eventually, 421(79.6%) survivors were contacted and reviewed by the supportive care team. After review, an additional 63/421 (15%) highly distressed survivors were referred to the multidisciplinary care team.

## FEASIBILITY – ACCESS TO COMMUNITY SERVICES

The supportive care team proposed community service referrals to 143/421 (34.0%) reviewed survivors, with 22 (5.2%) survivors receiving more than one referral (Table 6). Overall, the most common referral was to community

hospice (69/143, 48.3%), followed by cancer-specific community services (28/143, 19.6%), primary care physicians (27/143, 18.9%), and community facilities (18/143, 12.6%). Acceptance rates (Table 7) of these four

common referrals all exceeded 50%, with the highest acceptance rates for hospice (81.2%) and primary care physicians (81.5%). Attendance rates to accepted referral visits were the highest for hospice (79.3%) and

COMMUNITY SERVICE	PROPORTION OF HIGHLY DISTRESSED SURVIVORS WHO RECEIVED COMMUNITY REFERRALS BY THE SUPPORTIVE CARE TEAM, N (%)
Hospice	69 (13.0%)
Cancer-specific services <sup>a</sup>	28 (5.3%)
Primary care physicians	27 (5.1%)
Community facilities <sup>b</sup>	18 (3.4%)
Social services <sup>c</sup>	9 (1.7%)
Home-based care services	8 (1.5%)
Transport-related services	4 (0.8%)
Exercise programs	3 (0.6%)
Others <sup>d</sup>	6 (1.1%)

**Table 6** Type and frequency of community service referrals proposed to survivors reporting high distress (N = 529).

<sup>a</sup>Includes 365 Cancer Prevention Society, AIN Society, Breast Cancer Foundation, Children's Cancer Foundation, Singapore Cancer Society.

<sup>b</sup>Includes community day care facilities, community nursing, rehabilitative centres.

<sup>c</sup>Includes general, family, financial.

<sup>d</sup>includes eldercare, Ambulance Wish, Brave Charismatic, CGH Neighbours, private acupuncture, specialized disabled centre.

COMMUNITY SERVICE	ACCEPTANCE, N (%)	ATTENDANCE, N (%)
Hospice (n = 69)		
Referred/attended	58 (81.2%)	46 (79.3%)
Declined	11 (15.9%)	9 (15.5%)
Unknown/pending <sup>a</sup>	0 (0%)	3 (5.2%)
Cancer-specific services <sup>b</sup> (n = 28)		
Referred/attended	17 (60.7%)	11 (64.7%)
Declined	7 (25.0%)	1 (5.9%)
Unknown/pending <sup>a</sup>	4 (14.3%)	5 (29.4%)
Primary care physicians (n = 27)		
Referred/attended	22 (81.5%)	10 (45.5%)
Declined	3 (11.1%)	3 (13.6%)
Unknown/pending <sup>a</sup>	2 (7.4%)	9 (40.9%)
Community facilities <sup>c</sup> (n = 18)		
Referred/attended	12 (66.7%)	10 (83.3%)
Declined	6 (33.3%)	0 (0%)
Unknown/pending <sup>a</sup>	0 (0%)	2 (16.7%)

**Table 7** Acceptance rates of community service referrals and subsequent attendance rates to accepted referral visits.

<sup>a</sup>Refers to scheduled community visit that occurs after the study review period or the outcome of the community visit could not be traced.

<sup>b</sup>Includes 365 Cancer Prevention Society, AIN Society, Breast Cancer Foundation, Children's Cancer Foundation, Singapore Cancer Society.

<sup>c</sup>Includes community day care facilities, community nursing, rehabilitative centres.

community facilities (83.3%). A lower attendance rate for the remaining two services was accompanied by a higher proportion of referrals with visits planned after the study period or with untraceable outcomes.

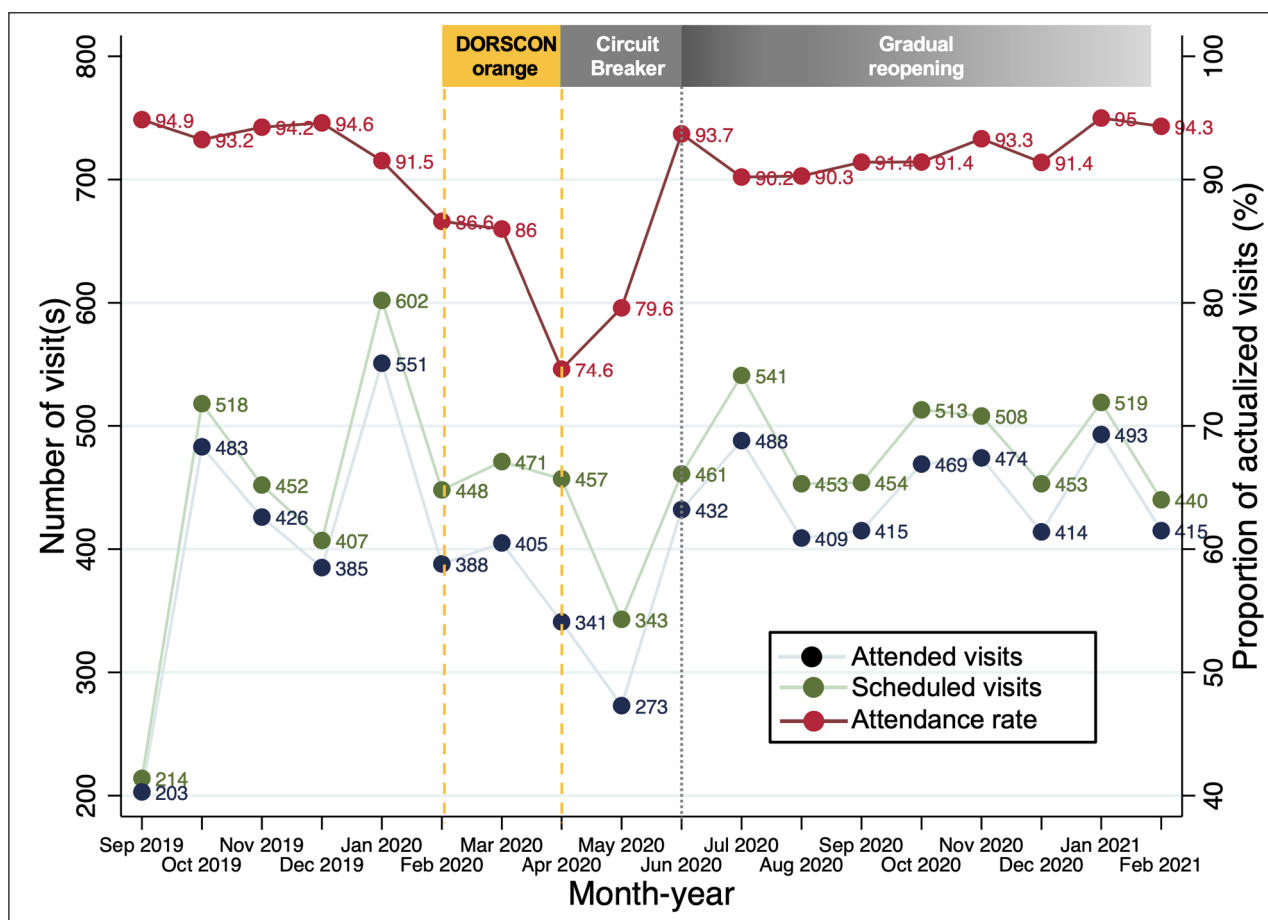
**IMPACT OF COVID-19**

Both scheduled and attended visits were negatively affected by COVID-19, with a trough corresponding to the lockdown (Figure 5). Model penetration was reduced between February to June 2020 as the number of clinic visits dropped to <400. Caseload recovered to the average monthly level of 500 visits after gradual reopening,

comparable with pre-COVID-19. Examining adherence to screening tool completion at different periods (Table 8), adherence rate decreased from 81.1% pre-COVID-19 to 64.6% during tightened measures, before returning to pre-COVID-19 levels (79.2%) after gradual reopening.

**DISCUSSION**

We presented results from the 18-month review of the ACCESS model which was successfully introduced to 1853 breast and gynaecological cancer survivors at



**Figure 5** Number of scheduled and attended visits, and proportion of attended visits over the 18-month implementation period when significant COVID-19 restrictions were imposed. (DORSCON = Disease Outbreak Response System Condition, with orange representing severe disease and tightened control measures; Circuit Breaker = lockdown).

OUTCOME	SIGNIFICANT COVID-19 EVENTS <sup>a</sup>			
	PRE-COVID-19 PERIOD	DORSCON ORANGE	CIRCUIT BREAKER	GRADUAL REOPENING
Attended visits, n (%)	2048 (93.4%)	793 (86.3%)	614 (76.8%)	4009 (92.3%)
Attended visits with screening tool completion, n (%)	1660 (81.1%)	512 (64.6%)	434 (70.7%)	3173 (79.2%)

**Table 8** Screening completion rates at attended medical oncologist visits stratified by periods of significant COVID-19 events.

<sup>a</sup> Defined by the following time periods: pre-COVID-19 lasted from September 2019 to end January 2020, DORSCON Orange with social distancing and mobility restriction measures lasted from February to end March 2020, Circuit breaker (lockdown) lasted from April to early June 2020, gradual reopening phase refers to the remaining time periods. Abbreviation: COVID-19, coronavirus 2019; DORSCON, Disease Outbreak Response System Condition.

an Asian ambulatory cancer centre in Singapore. Our findings demonstrated that a multidisciplinary care model like ACCESS, which employed routine screening with subsequent tiered care support using integrated care pathways, is feasible for incorporation into routine care. Additionally, our experience elucidated opportunities to improve referrals to community services and strategies to be inclusive of a diverse population in high-resource care settings to improve integrated cancer survivorship care provision.

Several studies evaluated the implementation of cancer supportive care models with distress screening following centre accreditation requirements and guidelines adherence [14, 17]. Our screening protocol demonstrated satisfactory feasibility after successfully screening the majority (78.6%) of eligible survivors despite COVID-19, close to the upper bound of the 12–86% range reported in the literature [39–41]. Long-term adherence was exemplified by high fidelity to screening despite multiple visits, an important indicator as survivors may only report high distress at repeated visits. We drew strength from two crucial strategies that could extend across settings. First, regular engagement with involved personnel facilitated an iterative feedback and improvement process. Workflows were streamlined and clinical efficiency improved with increased experience and additional training support [42, 43]. Second, a dedicated coordinator role served as a safety net to assist screening completion and guide survivors through the screening interface. This touchpoint reversed concerns of poor response rates [44, 45] as the majority (67.6%) of responses were completed by survivors independently. Collectively, an engaged workforce improved the model's feasibility and fidelity.

The ACCESS model identified 28.6% of screened survivors as highly distressed for supportive care team review, comparable to the 22–30% range reported in other outpatient cancer populations using DT cut-off score of 5–6 [46–49]. Screening allowed survivors to report emotional and practical problems, directly rectifying the current disproportionate focus on physical symptoms. Furthermore, our care team demonstrated high responsiveness to survivors' care needs. Specifically, multidisciplinary care meetings ensured holistic support for survivors with complex needs who would not have received a comparable level of care intensity under usual care. Additionally, the supportive care team proposed a diverse range of community service referrals to 34% of survivors reporting high distress, promoting access to community services. Educational activities conducted for community partners increased awareness of available community services that could benefit survivors in the cancer centre. Although the absolute numbers of referrals to some services were low, we observed >50% acceptance and attendance rates to referral visits. Consistent with local and international calls for shifting

survivorship care to community settings [21, 50, 51], the ACCESS model epitomized a systematic approach to engage community services from tertiary settings through mutually beneficial educational and outreach initiatives.

Our experience demonstrated the impact of integrating inclusivity into the design of supportive care models. Inclusivity at the screening stage is pivotal in ensuring that all survivors have an avenue to report care needs. Echoing the need for improved care accessibility in underserved populations, our results revealed the ACCESS model's inclusiveness of the multi-ethnic and multilingual Singapore population [19]. The ethnic distribution of our survivor cohort mirrored the distribution in the Singapore National Cancer Registry, suggesting a low likelihood that any ethnic group was systematically excluded [28]. Also, approximately 1 out of 4 survivors would have been deterred if non-English tools were unavailable. The availability of service coordinators' assistance for approximately 33% of responses obtained ensured that the model remained inclusive of survivors with poor literacy and technological knowledge. Adherence to screening did not differ by language choice and ethnicity, except for a 15% marginally lower completion rate among Malays than Chinese. This observation could be potentially explained by differences in perceived health status and health-seeking behaviour. Singapore-based studies revealed that Malays tend to report better self-rated health and lower adherence to cancer-related screening, reducing the perceived need for additional supportive services and screening for their cancer management [52, 53]. Nevertheless, the overall general positive results on inclusivity underscored efforts to develop screening resources in multiple languages, multiple modes, and ensuring cultural diversity in the workforce.

Strategies targeting inclusivity further allowed the model to be flexible and adapt to restrictions imposed by the COVID-19 pandemic. Notably, service coordinators switched promptly to administer DTPL via phone calls, achieving >50% screening completion rates even during tightened restrictions. However, survivors without phone access likely remained underserved during COVID-19. Therefore, a hybrid mode comprising electronic platforms, phone calls, and in-person encounters should be adopted or preserved to maintain inclusivity and robustness to external changes such as the pandemic.

There were several areas for improvement in our care model. We encountered a significant barrier with screening mechanics and poor integration with electronic medical records, consistent with other centres' experiences [54]. Extensive manpower utilized to support the manual components of screening incur significant opportunity costs that could potentially be reduced with digital solutions [55, 56]. Coordinators could have played a more active role as health coaches or care navigators

to improve outreach to underserved populations, such as the Malay group identified in our context, to further boost inclusivity [19, 57]. Also, the COVID-19 pandemic compromised the model's penetration due to deferred visits, failing to hit the expected caseload by almost 40% [58]. Lastly, community service referrals were impeded by a significant heterogeneity in referral workflows across community providers.

This study has several limitations. First, demographic and clinical characteristics of survivors who refused the model were not collected to reveal potential systematic differences. Second, the mode of screening tool completion served as a proxy for literacy as associated data were not routinely collected. Third, we could not ascertain how consistently screening results were discussed between oncologists and survivors. Fourth, we acknowledge potential limitations with the use of DT for screening, including the unknown optimal interval between repeated screening and unclear survivors' perceptions of high distress [59]. Notwithstanding, the DT is brief for routine administration. Furthermore, the overall high positive and negative predictive values of >80% corresponding to the cut-off score employed strongly justify its usage [34]. Lastly, the evaluation of the implementation process was mainly conducted from a health system's perspective. Thus, future work is underway to explore survivors' and their caregivers' experiences with the care model to supplement outcomes from this study. Additionally, the concurrent clinical trial as part of the hybrid study approach will elucidate the clinical impact of the model and facilitate subsequent economic evaluation to support long-term model adoption.

## LESSONS LEARNED

- Recognizing diverse care providers in integrated care provision, prospective adopters of a multidisciplinary supportive care model for cancer survivors in the outpatient setting should devote sufficient resources to engage the workforce continuously to sustain a positive implementation climate. Notably, a service coordinator role, preferably a person with health science-related background, should be carefully envisioned to complement the care team by providing general cancer survivors with navigation support or basic health coaching advice.
- A systematic approach should be adopted to facilitate access to community services and care providers from the tertiary settings. Foremost, mutually beneficial educational and outreach initiatives constitute a preliminary step to improve awareness of available community services and encourage collaboration opportunities. The derivation of decisive benefits from this interaction promotes mutualism to optimize the level of inter-organizational integration [60]. Community services should also be mapped to reported care needs in the integrated care pathways used in tertiary settings to encourage care coordination. A centralized referral system is further proposed to ensure care continuity through improved navigation and referral outcomes tracking.
- Key implementation strategies employed to ensure inclusivity of minority groups include developing screening resources in multiple languages, hybrid modes (electronic platforms, phone calls, and in-person encounters), and ensuring cultural diversity in the workforce. Further refinement of this approach should consider context-specific factors of the target setting and incorporate existing effective outreach strategies to minority groups.
- Inclusivity considerations provided additional positive impact as they allowed the care model to be flexible and remained robust to external changes such as the restrictions imposed by the COVID-19 pandemic. This flexibility is greatly enhanced by the inclusion of telehealth.

## CONCLUSION

Our multidisciplinary supportive care model designed with considerations of inclusivity and access to community services delivered integrated care to an ethnically diverse population. It demonstrated high fidelity, feasibility, and responsiveness to breast and gynaecological cancer survivors' care needs in the outpatient setting. Community service referrals were facilitated systematically through the supportive care team and a hybrid screening approach preserved inclusivity while allowing flexible adaptations during COVID-19. Complementing ongoing research to evaluate the effectiveness of the model on survivor-level outcomes such as quality of life and symptom burden, future research should explore telehealth's integration into supportive care models to streamline and promote integrated care. It would be useful to clarify how telehealth can further boost inclusivity and community engagement, providing access to multiple health disciplines and services through survivors' preferred modes dynamically. Furthermore, the potential for telehealth to transform a multidisciplinary care team into an interdisciplinary one will be of great interest to advance integrated care provision.

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## COMPETING INTERESTS

The authors have no competing interests to declare.

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