Consent for Systemic Anti-Cancer Therapy (SACT)

Guidance issued by the UK Chemotherapy Board
June 2018
Contents

1. Introduction
   1.1. Purpose of the guidance
   1.2. Systemic Anti-Cancer Therapy (SACT) and consent
   1.3. National report recommendations
   1.4. SACT consent forms: Current status

2. National SACT regimen-specific consent forms
   2.1. UK Chemotherapy Board recommendations
   2.2. Routine SACT regimens
   2.3. Clinical trials
   2.4. Chemo-radiation

3. Guidance for the introduction of the national SACT regimen-specific consent forms by individual chemotherapy providers
   3.1. Local Chemotherapy/SACT group
   3.2. Governance/Consent committee
   3.3. Local process for use of the forms

4. Guidance for the process of consent for SACT: Clinical use and completion of SACT regimen-specific consent forms
   4.1. Definition of consent
   4.2. Who can take consent?
   4.3. Who can give consent?
   4.4. When and where should consent for SACT be taken?
   4.5. How should consent for SACT be obtained and documented?
   4.6. Second consultations and confirmation of consent

5. Audit of consent procedures

6. Acknowledgments, consultation and support

7. References and bibliography

Appendix I

Appendix II
Template letter for governance/consent committees
1. Introduction

1.1. Purpose of the guidance

The purpose of this guidance is to:

- Describe the context relating to the requirement of consent for the administration of systemic anti-cancer therapy (SACT) and outline relevant recommendations from national reports.
- Introduce and recommend the use of SACT regimen-specific consent forms for all adult patients and launch the national library of forms.
- Provide guidance for all chemotherapy providers in relation to the local adoption and implementation of national SACT regimen-specific consent forms.
- Outline guidance relating to the process of providing information and obtaining consent from adults for treatment with SACT.
- Make recommendations for the audit of consent procedures.

1.2. Systemic Anti-Cancer Therapy (SACT) and consent

The use of systemic anti-cancer therapy (SACT) is increasing year on year, and the types of agents are growing with various new forms of treatment now available which may complement or replace conventional cytotoxic chemotherapy. Treatment with SACT is associated with complex risks with respect to administration and toxicity. Additionally, the risks and benefits of receiving these treatments will differ from patient to patient and, at times, this balance of risk, with respect to toxicity, will need to be carefully considered alongside any potential benefit in terms of survival or symptom control. Because of these issues, the procedure for consent for receipt of SACT involves almost unique uncertainties, and the process of obtaining consent requires considerable expertise and carries specific responsibilities.

Although written consent is not required in law, it is generally accepted that the prescription of SACT is best supported by written consent following a full discussion of the intended benefits and the associated risks. The signing of a consent form indicates that a process has taken place. It does not, necessarily, indicate that the patient has full comprehension of the treatment procedures, aims and complications. Completion of a consent form must, therefore, be complemented by contemporaneous records within the patient notes of discussions which have been held.

Several guidance documents have described best practice in the area of consent, with respect to law, ethics, training and experience required, and the need for documentation that consent has taken place. Following the Montgomery case in 2015 the law now requires a doctor to take ‘reasonable care to ensure that the patient is aware of any material risks involved in any recommended treatment, and of any reasonable alternative or variant treatments’. Renewed procedural guidance, based on these existing guidance documents, recent UK case law, and current perceived best practice is given in section 4 of this guidance.
1.3. National report recommendations

The National Chemotherapy Advisory Group (NCAG) published its report ‘Chemotherapy Services in England: Ensuring quality and safety’ in August 2009. The NCAG report was produced to address the significant concerns raised regarding the safety and quality of chemotherapy services in England in the 2008 National Confidential Enquiry into Patient Outcomes and Death (NCEPOD) and cancer peer review reports 2004-2007. The NCAG report made recommendations relating to the whole of the chemotherapy care pathway, i.e. from referral to an oncologist through to the end of treatment.

The main proposed actions within the NCAG report in relation to consent and information were that:

- ‘Standardised consent forms are used which include details of the toxicities discussed and which identify whether a patient has been provided with written information.’
- ‘A copy of the consent form should be given to the patient as well as one being filed in the patient’s case record.’
- ‘Standardised written information and other forms of information are given to patients, which are relevant to a particular chemotherapy regimen.’
- Chemotherapy service providers should ‘provide written information to patients about the chemotherapy they will be receiving, the likely side effects and whom they should contact if problems arise (including out of hours). Delivery of such information should be documented.’

The report recommended that ‘Consent should be obtained and recorded in detail in terms of the aims of treatment and both the common and serious side effects of treatment’. It also recommended that the quality of assessment and decision making should be assessed through audits of consent procedures coupled with patient experience surveys.

Consent is covered in some degree as part of the National Cancer Peer Review Programme in the Manual for Cancer Services: Chemotherapy Measures. There is a specific measure relating to the consent form (14-3S-132) and also checks prior to the prescription of the first cycle of chemotherapy (14-3S-127).

1.4. SACT consent forms: Current status

The Department of Health (DH) published guidance and four standard consent forms for use in the NHS in 2001. Although DH Consent Form 1 is appropriate for interventions where consciousness is not impaired, it lacks the structure to provide the detail that most cancer clinicians and ethicists would consider adequate to support and document the process of consent for SACT. The DH guidance was updated in 2009 but the gap between the available forms and the requirements for SACT remains.

Because of the limitations of the standard forms available, practice varies widely around the UK with some chemotherapy providers using or adapting generic DH consent forms and others developing more specific documentation that may or may not reflect the DH forms and which differ in content and style.
2. National SACT regimen-specific consent forms

2.1. UK Chemotherapy Board recommendations

The UK Chemotherapy Board is recommending the use of standardised SACT regimen-specific consent forms in the UK. This will support chemotherapy providers to adhere to best practice guidance, and address the recommendations and proposed actions in the NCAG report that are relevant to consent. Introduction and adoption of the national forms by all chemotherapy providers will confer the governance and quality benefits of standardised regimen-specific forms to all eligible patients.

SACT regimen-specific consent forms were developed as part of a project funded by the now disbanded South East London Cancer Network (SELCN). This library of forms has been maintained and updated by Guy’s and St. Thomas’ NHS Foundation Trust (GSTFT) oncology pharmacists and clinicians since 2010. There are over 200 forms available and in use within South East London. Since 2016, this resource, supported by funding from Cancer Research UK (CRUK), has been used to develop and launch the national library of standardised SACT regimen-specific consent forms which are available to all chemotherapy providers in the UK via the Cancer Research UK website (www.cancerresearchuk.org).

The national SACT regimen-specific forms include a minimum set of data that is necessary for patient’s information, and a minimum number of fields to be completed to support the process and identify the completion of responsibilities within the process. It is recognised that some organisations have specific requirements within their own consent procedures that may be outside the content of the forms recommended. Those organisations will be free to augment the recommended national forms, however, it is not envisaged that any content should be deleted locally.

2.2. Routine SACT regimens

National SACT regimen-specific consent forms will be made available for all oncology and haemato-oncology regimens that are in routine use within the UK for the treatment of adult patients with cancer. A generic SACT regimen-specific consent form template will also be available which may be used as an interim measure until all national forms are published. See appendix I for an example of a regimen-specific consent form.

2.3. Clinical trials

Many patients with cancer are treated within the setting of a clinical trial. Where SACT treatment is within a clinical trial, the consent for systemic therapy is no different and should follow this guidance.

Trial consent forms do not always contain specific details of the SACT regimen. Where this is the case, separate written consent should be obtained for participation in the trial and then either an existing regimen-specific or the generic regimen-specific form used for consent for the receipt of SACT. This duplication of consent is already embedded in practice in many organisations.
2.4. Chemo-radiation

These forms are designed specifically for the taking of consent for SACT. For instances where the SACT is given in combination with radiotherapy, the radiation therapy will need to be consented for separately.

3. Guidance for the introduction of the national SACT regimen-specific consent forms by individual chemotherapy providers

The process for the introduction and local adoption of national SACT regimen-specific consent forms may differ for every chemotherapy provider. Key aspects for each provider to consider in relation to the local introduction of the forms are described below.

3.1. Local Chemotherapy/SACT group

This guidance document and the national regimen-specific consent forms should be tabled for discussion by the local chemotherapy/SACT group, or equivalent. The group should benchmark their current SACT consent processes against this guidance. There should be consensus agreement to work in line with the guidance and adopt the use of the national forms, wherever possible. Ideally, they should nominate a lead person from their membership to take ownership for the introduction of the forms into local practice. They should develop and implement a process for the local use of the forms, to ensure appropriate adoption and observance of clinical governance, described in section 3.3 of this document.

3.2. Governance/Consent committee

A request to use the regimen-specific SACT forms should be submitted to the chemotherapy provider’s governance or consent committee, whichever is most appropriate to approach for the individual organisation. This submission can be supported by the SACT consent summary letter, a template for which is provided in appendix II, a copy of this guidance document and an example SACT regimen-specific consent form. The local process for the use of the forms could be sent with other supporting documentation, if this is available at the time of submission.

3.3. Local process for use of the forms

A standard operating procedure (SOP) or guidelines for use of the regimen-specific consent forms should be developed. This should describe the process for:
- Downloading and printing of the forms from the website, including appropriate version control;
- Storage of blank forms;
- Giving the patient a copy of completed forms;
- Filing of completed forms in the patient’s records.
4. **Guidance for the process of consent for SACT: Clinical use and completion of SACT regimen-specific consent forms**

4.1. **Definition of consent**

Consent to treatment is the principle that a person must give their permission before they receive any type of examination or medical treatment. For consent to be valid it must be voluntary and informed and the person consenting must have the capacity to make the decision. As described in section 1.3 of this guidance, although written consent is not required in law, it is generally accepted that the prescription of SACT is best supported by written consent following a full discussion of the intended benefits and the associated risks.

4.2. **Who can take consent?**

Consent should be obtained by a healthcare professional with sufficient training and experience who is also experienced in the management of the specific SACT regimen for which the patient will consent. The health professional must be aware of their legal, ethical and professional responsibilities with regards to consent, in addition to local institution policies and procedures.

It is expected that a competent healthcare professional will be one of the following:

- A consultant or associate specialist medical or clinical oncologist.
- A consultant or associate specialist clinical haematologist.
- A specialist registrar, clinical fellow or staff grade doctor in oncology or haematology who has had specific training in consent for systemic therapy.
- A nurse or pharmacist prescriber with specific training and documentation of training to prescribe the specific regimen for which the patient will consent.

4.3. **Who can give consent?**

This document does not seek to duplicate guidance given elsewhere, i.e. in the DH 2009 guidance and, its scope does not cover situations where the ability to give informed consent is lacking due to impaired understanding or consciousness. Guidance on this is summarised in the British Committee for Standards in Haematology (BCSH) guidelines on consent 2012 and those guidelines should be considered in conjunction with the principles within this document.

4.4. **When and where should consent for SACT be taken?**

Consent should be obtained and documented prior to the first cycle of a course of SACT using a defined number of cycles of the same regimen or a defined standard sequential treatment comprising defined numbers of cycles of more than one regimen.
Ideally there should be time (measured in days rather than hours) for the patient to reconsider their decision to undergo the treatment before the first cycle is commenced. It is acknowledged that within many cancer pathways, considerable discussion may have been held between a patient and their key worker or other healthcare professionals prior to a first consultation with a prescriber of SACT, however, full explanation of risks and benefits must still be undertaken during this consultation.

Consent should be taken within a consultation in which:

- The patient (and/or representative) has been introduced to the person taking consent and understands who they are.
- The person taking consent can demonstrate that they have sufficient knowledge of the patient’s individual circumstances (in particular co-morbidities) and details of their cancer (usually through a structured consultation and discussion of surgery and/or test results).
- There is sufficient privacy to discuss any issues that may arise.
- Any examinations have been undertaken with respect to dignity.
- The patient has had the opportunity to involve a relative, friend and/or carer.
- The patient has been able to request the presence of their key worker as defined in the Manual of Cancer Services.
- The patient and/or relative/friend/carer has had an opportunity to ask any questions that they may have.

Where English is not the first language of the patient translation facilities must be available to support the giving of information, answering of questions and taking of consent. It is advised that this should be through a professional translation service in preference to family or friends, other than in exceptional circumstances.

4.5. How should consent for SACT be obtained and documented?

The SACT regimen-specific form should be used as an aid to the discussions about treatment with SACT. Discussions should be supported by information leaflets for the specific regimen or constituent medicines, e.g. Cancer Research UK or Macmillan information sheets. The risks and benefits of the treatment, including response rates and alternative or variant options, should be discussed with the patient in terms that they can understand.

Key aspects of the discussion, which should be documented on the completed form are:

- The aims and intent of the treatment.
- The regimen and constituent medicines.
- The route of administration.
- What the treatment is likely to involve:
  - The schedule of administration and intended duration of treatment with the specific SACT regimen.
  - The location of administration (e.g. day-case unit).
  - Description of blood tests and additional tests (e.g. CT scan) and procedures (e.g. PICC line).
  - Outpatient and follow-up appointments.
- Short and long-term toxicities, and details of how these may be managed.
The common side-effects and life-threatening complications, even where these are rare.
  - The healthcare professional must take into account any co-morbidities the patient may have, and how these might impact any relevant toxicities.
- Effects on fertility, where relevant, must be discussed.
- The form must allow the consenting professional to confirm that advice on avoidance of pregnancy and conception has been provided where relevant.
- How and when to contact the hospital team if the patient has any problems or queries regarding their treatment.

The consenting healthcare professional must sign the form confirming that the discussions have taken place and that the patient has no further questions and wishes to proceed with treatment. They must ensure that all relevant sections of the form have been completed.

The interpreter should sign the form, where appropriate, confirming that they have interpreted the information to the patient to the best of their ability and in a way which they believe the patient can understand.

The patient should sign the form, once they have confirmed that they wish to proceed with treatment.

A copy of the completed form should be given to the patient with other appropriate information leaflets. The original signed consent form should be filed in the patient records.

**4.6. Second consultations and confirmation of consent**

The Manual of Cancer Services recommends that all patients should have the opportunity for a routine second consultation with a suitably trained and experienced healthcare professional prior to commencing treatment\(^8\) to ensure that they understand:

- Why the treatment has been offered.
- What alternatives could be offered.
- What the treatment involves.
- What the risks might be.
- What the likely benefits might be.
- That they can withdraw consent at any time.

The requirement for efficient delivery of cancer treatments and the need for urgent therapy should not be limited by the need for this second routine consultation. Hence, the second routine consultation can be performed by a competent chemotherapy nurse (or equivalent), prior to administration of the first cycle of chemotherapy.

Consent must be confirmed and documented on the consent form prior to the administration of the first cycle of SACT where the patient has signed the form in advance, i.e. most instances. This can be done by the treating nurse on the day of treatment. It is good practice to ensure ongoing consent throughout the planned course of treatment, although documentation on the specific form should only be required prior to the first cycle.
5. Audit of consent procedures

Periodic audit of consent procedures for SACT are recommended. As a minimum, the audit should include the following standards where 100% compliance should be seen:

- Consent forms are available for review.
- The name and job title of the responsible healthcare professional are documented.
- The treatment intent and intended benefits are documented.
- The common toxicities of the SACT are recorded.
- The form has been signed and dated by the healthcare professional and the patient.

6. Acknowledgements, consultation and support

This guidance document was initially developed by a working group of the National Chemotherapy Implementation Group (NCIG), which was dissolved in 2012 with the reorganisation of the Department of Health and the creation of NHS England. The UK Chemotherapy Board acknowledges the work that was done by our predecessors in developing this guidance.

The content of this guidance document has been reviewed and ratified by the UK Chemotherapy Board which has representation from the following National bodies:

- The Association of Cancer Physicians (ACP).
- The Royal College of Radiologists (RCR).
- The Royal College of Physicians (RCP).
- The Royal College of Pathologists (RCPPath).
- The UK Oncology Nursing Society (UKONS).
- The British Oncology Pharmacy Association (BOPA).

All representatives have had opportunity to comment and make suggestions on the content of the document and the template regimen-specific consent form.

Cancer Research UK has awarded a grant to Guy’s and St. Thomas’ NHS Foundation Trust to host and deliver the National SACT regimen-specific consent form project. They are supporting production and hosting of the forms on the CRUK website.
7. References and bibliography

1. Reference guide to consent for examination or treatment, Department of Health, 2009.
6. For better, or worse? National Confidential Enquiry into Patient Outcome and Death, November 2008.

![Consent Form Example](Image)
STATEMENT OF HEALTH PROFESSIONAL (continued)

SIGNIFICANT, UNAVOIDABLE OR FREQUENTLY OCCURRING RISKS

COMMON SIDE EFFECTS:
More than 10 in every 100 (1-10%) people have one or more of the side effects listed:

- Diarrhoea, feeling sick (nausea) and being sick (vomiting), sore mouth and ulcers, taste changes, anaemia (low number of red blood cells), hair loss, bruising or bleeding, tiredness and feeling weak (fatigue), numbness or tingling in the hands and feet which may be temporary or persistent, skin changes, aching or pain in joints and muscles, headaches, low blood pressure during treatment, and changes in the way the kidneys work.

- An increased risk of getting an infection from a drop in white blood cells - it is harder to fight infections and you can become very ill.

If you have a severe infection this can be life-threatening. Contact your doctor or hospital straight away if:

- your temperature goes over 37.5°C (99.5°F) or over 38°C (100.4°F), depending on the advice given by your chemotherapy team
- you suddenly feel unwell (even with a normal temperature)

OTHER RISKS:

- Paclitaxel may leak outside of the vein while it is being given; this is called extravasation. If this happens when you’re having paclitaxel, it can damage the tissue around the vein. Tell the nurse straight away if you have any stinging, pain, redness or swelling around the vein. Extravasation is not common but if it happens it’s important that it’s dealt with quickly.

- Paclitaxel and carboplatin may cause an allergic reaction while being given. You will have medicines before your treatment to help prevent this.

- Potential side-effects with the anti-sickness medication may include: constipation, headaches, indigestion, difficulty sleeping, and agitation.

- Cancer can increase your risk or developing a blood clot (thrombosis), and having treatment with anti-cancer medicines may increase this risk further. A blood clot may cause pain, redness and swelling in a leg, or breathlessness and chest pain - you must tell your doctor straight away if you have any of these symptoms.

- Some anti-cancer medicines can damage women’s ovaries and men’s sperm. This may lead to infertility in men and women and/or early menopause in women.

- Some anti-cancer medicines may damage the development of a baby in the womb. It is important not to become pregnant or father a child while you are having treatment and for at least 6 months afterwards. It is important to use effective contraception during and for at least 6 months after treatment. You can talk to your doctor or nurse about this.

- Very rarely complications of treatment with anti-cancer medicines can be life-threatening or even result in death. The risks are different for every individual. You can talk to your doctor or nurse about what this means for you.

---

TO BE RETAINED IN PATIENT NOTES
Prepared by: René Chauhan
Checked by: Pharmacist: SAMPLE
Checked by Consultant: SAMPLE

Date of issue and version: Jan-27, Version SAMPLE
Reviewed: Jan-20
Approved by: Janine Mansi (UK Chemotherapy Board)
Check www.cancerresearchuk.org for latest version

Page 2 of 5

Consent for Systemic Anti-Cancer Therapy (SACT) 13
STATEMENT OF HEALTH PROFESSIONAL (continued)

ANY OTHER RISKS:

☐ I have discussed what the treatment is likely to involve (including inpatient / outpatient treatment, timing of the treatment, blood and any additional tests, follow-up appointments etc) and location.
☐ I have discussed the intended benefits and risks of any available alternative treatments (including no treatment) and any particular concerns of this patient.

THE FOLLOWING LEAFLET/TAPE HAS BEEN PROVIDED:

☐ Information leaflet for carboplatin-paclitaxel chemotherapy and/or for the individual drugs (carboplatin and paclitaxel).
☐ 24-hour chemotherapy service contact details

Signed:

Name (PRINT): _____________________________
Job title: _____________________________

Date: _____________________________

☐ Other, please state:

Signed:

Name (PRINT): _____________________________
Job title: _____________________________

Date: _____________________________

STATEMENT OF INTERPRETER (where appropriate)

INTERPRETER BOOKING REFERENCE (if applicable):

I have interpreted the information above to the patient to the best of my ability and in a way in which I believe s/he can understand.

Signed:

Name (PRINT): _____________________________
Job title: _____________________________

Date: _____________________________

TO BE RETAINED IN PATIENT NOTES
Prepared by: Rena Chauhan
Checked by: Pharmacists SAMPLE
Checked by: Consultant SAMPLE

Date of issue and version: Jan-17, Version SAMPLE
Review date: Jan-20
Approved by: Janine Marsh (UK Chemotherapy Board)
Check www.cancerresearchuk.org for latest version

Page 3 of 5
STATEMENT OF PATIENT

Please read this form carefully. If your treatment has been planned in advance, you should already have your own copy of the form which describes the benefits and risks of the proposed treatment. If not, you will be offered a copy now. If you have any further questions, do ask – we are here to help you. You have the right to change your mind at any time, including after you have signed this form.

I agree to the procedure and course of treatment described on this form.

I understand that you cannot give me a guarantee that a particular person will perform the procedure. The person will, however, have appropriate training and experience.

I understand that any procedure in addition to those described on this form will only be carried out if it is necessary to save my life or to prevent serious harm to my health.

I have been told about additional procedures which may become necessary during my treatment. I have listed below any procedures which I do not wish to be carried out without further discussion:

Patient's signature: ___________________________ Date: ____________
Name (PRINT): ____________________________________________

A witness should sign below if the patient is unable to sign but has indicated his or her consent. Young people/children may also like a parent to sign here (see notes).
Parent's/Witness's signature: __________________ Date: ____________
Name (PRINT): ____________________________________________

COPY ACCEPTED BY PATIENT: YES / NO
(please circle)

CONFIRMATION OF CONSENT
(health professional to complete when the patient attends for treatment, if the patient has signed the form in advance)

On behalf of the team treating the patient, I have confirmed with the patient that s/he has no further questions and wishes the procedure to go ahead.
Signed: ____________________________________________
Date: ____________
Name (PRINT): ____________________________________________
Job title: ____________________________________________

IMPORTANT NOTES: (tick if applicable)
☐ See also advance decision to refuse treatment
☐ Patient has withdrawn consent
  (ask patient to sign/date here)
Signed: ____________________________________________
Date: ____________

FURTHER INFORMATION FOR PATIENTS

CONTACT DETAILS (if patient wishes to discuss options later):

Contact your hospital team if you have any questions about cancer and treatment.
Cancer Research UK can also help answer your questions about cancer and treatment. If you want to talk in confidence, call our Information nurses on freephone 0800 800 4040, Monday to Friday, 9am to 5pm.
Alternatively visit www.cruk.org for more information.

These forms have been produced by Guy’s and St Thomas’ NHS Foundation Trust as part of a national project to support clinicians in ensuring all patients are fully informed when consenting to SACT. The project is supported by Cancer Research UK. This does not mean you are taking part in a clinical trial.

TO BE RETAINED IN PATIENT NOTES
Prepared by: Rena Chauhan
Checked by Consultant: SAMPLE
Date of issue and version: Jan-17; Version SAMPLE;
Review date: Jan-20
Approved by: Janine Mansi (UK Chemotherapy Board)
Check www.cancerresearchuk.org for latest version

Page 4 of 5
GUIDANCE FOR HEALTH PROFESSIONALS
(to be read in conjunction with the hospital’s consent policy)

WHAT A CONSENT FORM IS FOR
This form documents the patient’s agreement to go ahead with the investigation or treatment you have proposed. It is not a legal waiver – if patients, for example, do not receive enough information on which to base their decision, then the consent may not be valid, even though the form has been signed. Patients are also entitled to change their mind after signing the form, if they retain capacity to do so. The form should act as an aide-memoir to health professionals and patients, by providing a checklist of the kind of information patients should be offered, and by enabling the patient to have a written record of the main points discussed. In no way, however, should the written information provided for the patient be regarded as a substitute for face-to-face discussions with the patient.

THE LAW ON CONSENT
See the Department of Health’s Reference guide to consent for examination or treatment 2nd Edition for a comprehensive summary of the law on consent (also available at www.doh.gov.uk)

WHO CAN GIVE CONSENT
Everyone aged 16 or over is presumed to have the capacity to give consent for themselves, unless the opposite is demonstrated. If a child under the age of 10 has “sufficient understanding and intelligence to enable him or her to understand fully what is proposed”, then he or she will have capacity to give consent for himself or herself. Young people aged 16 and 17, and younger children with capacity, may therefore sign this form for themselves. Just like an adult patient, consent cannot be obtained for a child who is not able to give consent for himself or herself, unless someone with parental responsibility may do so on their behalf and a separate form is available for this purpose. Even where a child is able to give consent for themselves, you should always involve those with parental responsibility in the child’s care, unless the child specifically asks you not to do so. If a patient has the capacity to give consent but is physically unable to sign a form, you should complete this form as usual, and ask an independent witness to confirm that the patient has given consent orally or non-verbally.

WHEN NOT TO USE THIS FORM
If the patient is 16 or over and lacks the capacity to give consent, you should use an alternative form (form for adults who lack the capacity to consent to investigation or treatment) instead of this form. A patient lacks capacity if they have an impairment of the mind or brain or disturbance affecting the way their mind or brain works and they cannot:
- understand information about the decision to be made
- retain that information in their mind
- use or weigh this information as a part of their decision making process, or
- communicate their decision (by talking, using sign language or any other means)

You should always take all reasonable steps (for example involving more specialist colleagues) to support a patient in making their own decision, before concluding that they are unable to do so.

Relatives cannot be asked to sign a form on behalf of an adult who lacks capacity to consent for themselves, unless they have been given the authority to do so under a Lasting Power of Attorney or as a court deputy.

INFORMATION
Information about what the treatment will involve, its benefits and risks (including side-effects and complications) and the alternatives to the particular procedure proposed, is crucial for patients when making up their minds. The courts have stated that patients should be told about “significant risks which would affect the judgement of a reasonable patient”. “Significant” has not been legally defined, but the GMC requires doctors to tell patients about “significant, unavoidable or frequently occurring” risks. In addition if patients make clear they have particular concerns about certain kinds of risk, you should make sure they are informed about these risks, even if they are very small or rare. You should always answer questions honestly. Sometimes, patients may make it clear that they do not want to have any information about the options, but want you to decide on their behalf. In such circumstances, you should do your best to ensure that the patient receives at least very basic information about what is proposed. Where information is refused, you should document this on the consent form or in the patient’s notes.

REFERENCES
1. Summary of Product Characteristics (SPCs) for individual drugs: https://www.medicines.org.uk/emc
4. Guy’s and St. Thomas’ NHS Foundation Trust, Chemotherapy consent forms.

TO BE RETAINED IN PATIENT NOTES
Prepared by: Rena Chauhan
Checked by Pharmacist: SAMPLE
Checked by Consultant: SAMPLE
Date of issue and version: Jan-17, Version SAMPLE;
Review date: Jan-20
Approved by: Janine Mansi (UK Chemotherapy Board)
Check www.cancerresearchuk.org for latest version

Page 5 of 5
Appendix II – Template letter for governance/consent committees

Name and address of provider organisation

Date

To: Chair of Governance/Consent committee

Dear Chair name,

Re: Systemic Anti-Cancer Therapy (SACT) regimen-specific consent forms

The UK Chemotherapy Board has issued guidance relating to consent for SACT and has recommended the use of standardised SACT regimen-specific consent forms in the UK. To support this recommendation a national library of standardised SACT regimen-specific consent forms is being made available to all chemotherapy providers in the UK via a website. This is to support chemotherapy providers to adhere to best practice guidance, and address the recommendations and proposed actions in the National Chemotherapy Action Group (NCAG) 2009 report that are relevant to consent.

The name of Chemotherapy/SACT group have considered this recommendation and benchmarked our existing SACT consent practice against the national guidance. Our existing practice is brief description of existing practice, e.g. use of Trust/DH generic consent forms. Following this exercise the group have decided that they wish to adopt and implement use of the national SACT regimen-specific consent forms into local practice for the benefits of our service and ultimately our patients. To support this we have outlined a proposed local process for use of the forms, which is enclosed with this submission.

I am writing to submit a proposal to use the national SACT regimen-specific consent forms at name of provider organisation in preference to our existing practice.

Yours sincerely,

Chair of Chemotherapy/SACT group