What is a clinical trial?
An introduction for patients and families
Rona Passmore, Scotland:

I would absolutely recommend women to take part in trials if they feel it is right for them. Certainly I was very keen to take part in it because I want to try and increase the different drugs that are available for use with ovarian cancer. Without participation in trials, new drugs will not be tested and progress will not be made. Moving forward and increasing survival time is so important. My trial ran alongside my standard chemo initially, then carried on for a year. It took little time and was of no inconvenience to myself or my family. I worked full time throughout. I would encourage other women, if it’s right for them, to take part and help map a better outlook for those diagnosed with ovarian cancer.

Be positive.

Introduction

Life expectancy in Europe in the first decade of this century has continued to rise. We owe this fortunate development to various factors, not least the enormous progress made in the field of medicine. Today, many diseases are either curable or can at least be treated to enable prolong life expectancy. Clinical trials are pivotal in enabling this progress as they provide the scientific background for new medications, new treatment strategies and innovative diagnostic procedures.

Modern treatments are licensed by the European Medicines Agency (EMA) if they are proven to be more effective than the current standard of care and are monitored very closely in each country to ensure they are safe.

Before they can be used in clinical practice, they go through many years of development to ensure that they are safe and effective. On average it takes at least 10 years and more than 1 billion euros to develop a new drug.

This brochure is intended to give you comprehensive information about clinical trials in gynaecological cancers. It is designed to help you consider whether to participate in a clinical trial if you have been approached about taking part, and how you can find out more about participating in a trial.

Every person participating in a study does so on a completely voluntary basis and can also opt out of the study at any time without giving a reason.
What is a clinical trial?

Clinical trials, sometimes called studies, are a form of research which involves people. They are the final step in a long process that begins with research in a laboratory, and there are typically three phases in every clinical trial:

Three phases of a clinical trial

1. **In Phase I**, the new treatment is usually given at very low doses in the first instance, because the aim of this phase is primarily to examine how it affects the human body, to find a safe dose and to find how it should be delivered, although in cancer early signs of activity are hoped for. A Phase I trial is usually fairly small, typically involving 15-30 patients.

2. **In Phase II**, the treatment’s effectiveness is studied as well as its side-effects. Phase II trials involve a larger number of patients than Phase 1, but typically it will be less than 100.

3. **In Phase III**, the safety and effectiveness of the new treatment is usually tested against an existing optimum treatment or recognised standard of care. In a Phase III trial, patients are randomly allocated to the various treatment arms, in a process similar to throwing a dice, known as “randomisation.” The study group is given the new treatment or procedure to be tested; the control group is given an existing optimum treatment which may or not include a placebo. It is important to understand that patients cannot choose which group they will be allocated to, and that they may not receive the experimental treatment in the trial. Some may receive an inactive drug called a placebo, and this will be explained to you in the information sheet.

   A Phase III trial, which typically involves hundreds if not thousands of patients, is a prerequisite for official approval of a new treatment in Europe as this is the only way to evaluate the effectiveness of the new treatment compared to existing treatments.

   It should be noted that randomisation, typically associated with Phase III trials, is also increasingly used in Phase II trials.
CLINICAL TRIALS

Who conducts clinical trials?

Researchers conduct clinical trials in different settings. Many clinical trials are done at cancer centers because the facilities available are usually the most advanced. The research team that conducts a clinical trial can include research scientists, doctors, nurses, social workers, dietitians, and other healthcare professionals.

How are clinical trials approved?

The prerequisite for conducting any clinical trial is a detailed study protocol, namely a predefined written description of the intention and purpose, possible side-effects and the precise course of the trial, rights and duties of patients and plans of the analysis to be done in order to find out the results. An Ethics Committee of independent experts and lay people examine the trial protocol and decide whether, on the basis of the latest medical knowledge, it is purposeful and ethically justifiable to conduct the study. The Ethics Committee also examine whether the doctors, and institutions, who conduct these studies have the necessary knowledge and structures to administer such treatment.

Any change in the protocol, no matter how small, must be discussed with the person holding overall responsibility for the study and submitted for ethical approval. This high standard ensures that the safety and confidentiality of patients participating in a trial to develop new treatments are always the guiding principles of the trial.

Every patient participating in a clinical trial must be given comprehensive oral and written information before entering the trial, and sufficient time to consider it.

Why enter a clinical trial?

The fundamental principle of any trial is the safety and well-being of the trial participants. This takes priority over the interests of scientific research.

There are numerous good reasons for participating in a clinical trial. For patients it is comforting to know that as trial participants they are acquiring access to new and possibly better medications that in many cases would only be available to the general public several years later. In addition, being able to actively contribute to the progress of medicine and improvement of knowledge that will benefit future patients is a positive motivation.
Benefits of participating in a clinical trial

Another important advantage is the close monitoring of trial participants. Patients participating in clinical trials receive the best possible therapy available. They undergo particularly intense and meticulous medical examinations, and their treatment progress is precisely monitored. In the preparatory phase of the trial the dates for their medical examinations are set in the trial protocol, usually for several years in advance. At the beginning of the trial these patient examinations are scheduled at shorter intervals; in the first three years they are commonly conducted every three months and then every six months. Thereafter, patients are usually seen once a year.

This ongoing patient monitoring not only brings important findings in terms of scientific work, but also helps your doctors notice any decline in your health or a relapse at any early date so that the necessary steps can be taken quickly.

In most clinical studies the doctor responsible for the study and your care collects, analyses and checks not only the important health data obtained in the study, but also many other data. The advantage is that side-effects of treatment and changes in your health can be quickly recognised and treated. Furthermore, a clinical trial involves medical specialists from a wide range of disciplines, who are specialised in the management of the particular disease. This ensures optimum treatment whichever arm of the trial the patient is on.

Participation in clinical trials is always voluntary and always requires your written consent. Patients should understand what the trial’s goals and its special features are and what risks, if any, it entails. It should also be understood that new treatments that are being tested are not always better than, or even as good as, existing treatments and that they may have unexpected side effects.

Clinically trials offer hope for many people and an opportunity to help researchers find better treatments for others in the future.

Are you thinking about participating in a clinical trial?

Here are some of the questions you may wish to ask your oncologist:

- Am I eligible for a clinical trial?
- What is the purpose of the trial?
- What are the benefits?
- What are the side effects and risks?
- How many people are being recruited to the trial?
- What will it involve and how long will it last?
- How often will I have to visit the clinic?
- Will I be told the results?
- How can I find out what clinical trials are being offered?
- What kind of therapies, procedures and/or tests will I have during the trial?
- How will the trial affect my current treatment plan and affect my daily life?
- Will I be taken off my current treatment or will I continue to take my regular medications while in the trial?
- If the person in charge of my medical care during the trial is different from the doctor that put me on my current care, what will be the communication/interaction between the two?
- Can I talk to other people in the study?
- What is the procedure for opting out of this study once I have started it?
- Will there be costs to me if I participate?
- Will my insurance cover the costs?
- What are my responsibilities if I participate?
- How long will I be in the study?

Gynaecological cancer patients should not hesitate to volunteer for clinical trials because they have led to numerous benefits for patients throughout Europe, however, we recommend that they speak with their treatment team so that they are fully informed about what is involved before making a decision.
Clinical trials are research studies designed to evaluate the safety and effectiveness of new drugs or other types of therapies. This brochure has been specifically written as an educational tool for patients and their families providing an insight into how clinical trials work. It is a resource for those who may consider volunteering to participate in a clinical trial on the recommendation of their doctor.

ENGAGe is working with health professionals, policy makers and patients themselves to better understand the role of patients in providing input into clinical trial design and to achieve a better understanding of current barriers to participation in research.

Patients have a unique knowledge about their disease, treatment and quality of life. These insights are of great value to researchers and policymakers when priorities need to be set.

How to participate in a clinical trial?

Several ways are open to patients wishing to enquire about a clinical trial.

Some countries have national websites. The easiest is to ask your oncologist. There is also information about clinical trials on the following sites:

- European Union Clinical Trials Register: http://tinyurl.com/khevg4
- ESGO ENGOT: www.esgo.org/engot/pages/ENGOTrials.aspx
- Target Ovarian Cancer: http://clinicaltrials.targetovariancancer.org.uk

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Clinical trials are research studies designed to evaluate the safety and effectiveness of new drugs or other types of therapies.

Established by ESGO in 2012, the European Network of Gynaecological Cancer Advocacy Groups (ENGAGe) is a network of European patient advocacy groups representing all gynaecological cancers including ovarian, endometrial, cervical, vulvar and rarer gynaecological cancers under the umbrella of ESGO.

There are wide variations in patient care across Europe, including patients not being adequately informed about gynaecological cancers and their management. There are also issues, in relation to psychosocial support throughout the patient’s journey which may be less than ideal.

The objectives of ENGAGe are to:

- Facilitate the development of national gynaecological cancer patient groups in Europe and facilitate networking and collaboration between them.
- Disseminate information and share best practice to empower patient groups and improve the quality of care across Europe.
- To increase patient representation in ESGO activities by education on current research and health policy.
- To advocate patient care policies, practices and access to appropriate care at both national and European levels.
- To educate patient groups, health professionals, the public and health decision makers.
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