Patient involvement reports – Why they matter

Today you will see our patient involvement report on successful patient involvement, make sure you have a read.

Patients Involved | Case Report Patient Advocacy in HIV DUET phase III trials

Clinical Development Phase I , II & III

The pharmaceutical developer Tibotec (now Janssen Therapeutics) certain medicines or classes of medicines is more common with designed the DUET 1 & 2 studies in 2005. The DUET Phase III trials treatment experienced patients who therefore need novel or more involved the concurrent use of TMC125 (etravirine) and TMC114 complex regimens to control virus reproduction in the body. (darunavir) in a HIV treatment experienced population. The unique feature of the trial was that both compounds used had not been licensed at the time of use (2008). This was the first occasion that compounds. Standard procedure is to use a single new compound two yet unlicensed compounds were used in a trial in a treatment experienced setting, albeit only in one arm, while the other arm of the trial remained placebo-controlled.

requires a relatively rigorous regime of antiretroviral medication Compassionate use of the novel treatment regime through the trial (ART) for the patients in order to avoid resistance. Resistance to was advocated for.

Non-clinical Development

The patient community played a key role in achieving that -for the first time -a trial involved the concurrent use of two unregistered in a trial.

The objective of this intervention of the patient community was to make sure that a potent novel combination of ART is available HIV infection is a yet incurable but manageable disease that as salvage therapy for heavily treatment experienced patients.

Post-approval Life-cycle Management & Pharmacovigilance

Benefits

Research & Discovery

Consultation between the patient community and the pharmaceutical developer matured and evolved during this process significantly. The patient's organisations (POs) involved could successfully demonstrate to the industry and the regulators that the knowledge and experience of the patient community can yield substantial input into the development process. The innovative approach of the community infused the development process with a certain degree of 'courage' to go apply unconventional strategies when Providing compassionate use of novel compounds to patients preliminary results from previous trials are convincing enough (both new compounds were already known to be safe and well tolerable at the time).

This new approach led to lasting results and trust between the stakeholders involved. The collaboration of two POs from both sides Substantial advocacy (political) input was required from the patient of the Atlantic entered a new, more intensive phase, thus allowing community to convince the developer (and in turn FDA, EMA and exchange of experience across the communities of people living NCA) of this new strategy. with HIV. The DUET study resulted in overcoming accumulated MDR for thousands of heavily pretreated patients.

HIV infection is a yet incurable but manageable disease that requires a relatively rigorous regime of antiretroviral medication (ART) for the patients in order to avoid resistance. Resistance to However, one main challenge remained that the pharmaceutical certain medicines or classes of medicines is more common with treatment experienced patients who therefore need novel or more complex regimens to control virus reproduction in the body.

The patient community played a key role in achieving that -for the first time -a trial involved the concurrent use of two unregistered compounds. Standard procedure is to use a single new compound in a ṫrial.

The objective of this intervention of the patient community was to make sure that a potent novel combination of ART is available as salvage therapy for heavily treatment experienced patients. Compassionate use of the novel treatment regime through the trial was advocated for.

Who was involved?

EUPATI

www.eupati.eu This work is lief

European Patients' Academy on Therapeutic Innovation

Tibotec, AIDS Treatment Activists Coalition Drug Development Committee (USA), European Community Advisory Board (ECAB) of the European AIDS Treatment Group (EATG)

Level of patient expertise

- Patients with personal disease experience. - Expert patients / patient advocates with good expertise on

disease, but little R&D experience.

- Expert patients / patient advocates with good expertise on disease and good R&D experience.

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with reduced treatment options was and remains a challenge. The participation in clinical trials is an effective tool for patients to access new medicines.

The use of two experimental compounds was not common practice.

An important meeting was held with ATAC-DDC, EATG/ECAB and the pharmaceutical company in Antwerp in 2005. The specific objective of the meeting was to convince the company of the usefulness of and need for a new approach to help patients in need. company decided to design the trial with one placebo-controlled arm, meaning that 50% of the patients received placebo + one investigational compound rather than both new medicines.

earnings

The involvement of patient organisations and expert patients in pharmaceutical development is no longer unique. However, new strategies and uninterrupted work, complete with continuous selfeducation and rigorous knowledge of the field by the community are needed in order to navigate the complex setting of medicine development and research.

More intensive interaction with regulators is required to leverage the political objectives and pressure that POs want to exert to achieve their objectives; in this case the availability of new treatment options. Despite all efforts, the POs could only achieve a partial result: a placebo-controlled arm remained part of the trials concerned. Improvements in this area could, however, be achieved in later study designs developed with patient involvement.

Even better coordination between POs and a more regular exchange of experience within and outside a specific disease area should improve the effectiveness and efficiency of patient involvement in research

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A screenshot of a sample Patients Involved Case Report

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In the coming weeks we shall share patient involvement reports on trials and projects with successful patient involvement. Be sure to check back each week to see the new

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story. You will see that these case studies are from a wide range of diseases areas, and that patient advocates and patient experts were involved in the lots of different stages.

The definition of success is also not always entirely clearcut. In some cases, for example in the field of HIV/AIDS where I come from, success was sometimes stopping a trial, or just making sure that one or the other arm of a clinical study was modified to meet the patients' real-life needs more.

In other cases the end result was the creation of a patient advisory group that could effectively intervene and provide input into clinical trials.

One thing is common in all of them: They bear witness to the countless possibilities and opportunities that are there for patients to be involved, even if it takes a lot of time and effort! Some groups or disease areas might be more ahead, others are just stretching their wings in the world of patient activism. This collection of good practices, which does not stop short of admitting mistakes or failures as well, can be seen as both a motivational tool and some handy help for patient advocates.

Do you have a story you'd like to share? Contact the content team with a description of your case of patient involvement, making sure to include all required information so that it can be published. Help us grow our library of patients involvement reports!

Information required for patient involvement reports:

Partners Involved

Type of patient (advocates) involved (Patients with personal disease experience, and/or Expert patient / patient advocate with good expertise on disease, but little R&D experience, and/or Expert patient / patient advocate with good expertise on disease and good R&D experience, and/or, Other, describe

here:)

Description of the case (how were patients involved in the R&D project? What was the objective? (200 words max)

Benefits (how has this collaboration improved R&D process(es) and the R&D outcome(s) or triggered R&D organisational change) (150 words max)

Challenges and barriers (and how you have overcome them, or which ones were unresolved) (150 words max)

Discussion and learnings for you and EUPATI (what would you do differently next time, what are external factors that should change) (150 words max)

What phase of development did the involvement take place? (choose from the diagram below)



When does it happen?

Send your story to content@eupati.eu

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