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1997 Prof A. Potemkowski establishes Individual Specialist Neurological Practice. 1999 Prof A. Potemkowski signs a contract with NHS. 2000 Prof A. Potemkowski conducts first SM clinical trials. 2004 Dr M. Ratajczak and dr A. Ratajczak join Prof A. Potemkowski research team. 2005 Dr. J. Kolenda joins the research team. Dr M. Ratajczak establishes the Individual Doctor's Practice. 2006 The establishment of Memory Disorder Centre. 2009 The establishment of EuroMedis. Transfer of the NHS contract to EuroMedis. Extension of the NHS contract. Expanding the scope of Clinical Trials. Adding Specialist Clinics to our offer. 2014 Relocation to the new head office-further development of the Centre. 2021 Opening a 24-hour diagnostic department.

HISTORY



LOCATION

- The nearest airport Goleniów port serves domestic and international flights (UK, Scandinavia). The transfer to the airport takes about 40 minutes from Szczecin.
- Airport Berlin Schoenefeld 2 hours by car from Szczecin.







OUR MISSION

We believe that by providing health services focused on the needs of the patient and his family, we improve the quality of life, and because of our commitment to the development of new ideas and technologies we contribute to overcoming the limitations of current medicine.







OUR VISION

We want our services to be widely available, comprehensive and improve the quality of life so that everyone can enjoy the health.





ORGANIZATION









EuroMedis O



Over 50 specialists, 2 600 m2 of space in two locations.









DIAGNOSTICS

EuroMedis Medical Centre provides own diagnostics facilities.

Sampling for laboratory analysis, Ultrasonography, EEG, ECG with Holter monitoring and ergospirometry are all provided onsite.

Patients are also provided with a fully equipped ophthalmological laboratory where, for instance, perimetry is performed.

USG

EEG

Doppler USG

> **RR** Holter

ECG

ECG

Holter

Perimetry

Ergospirometry





DIAGNOSTICS

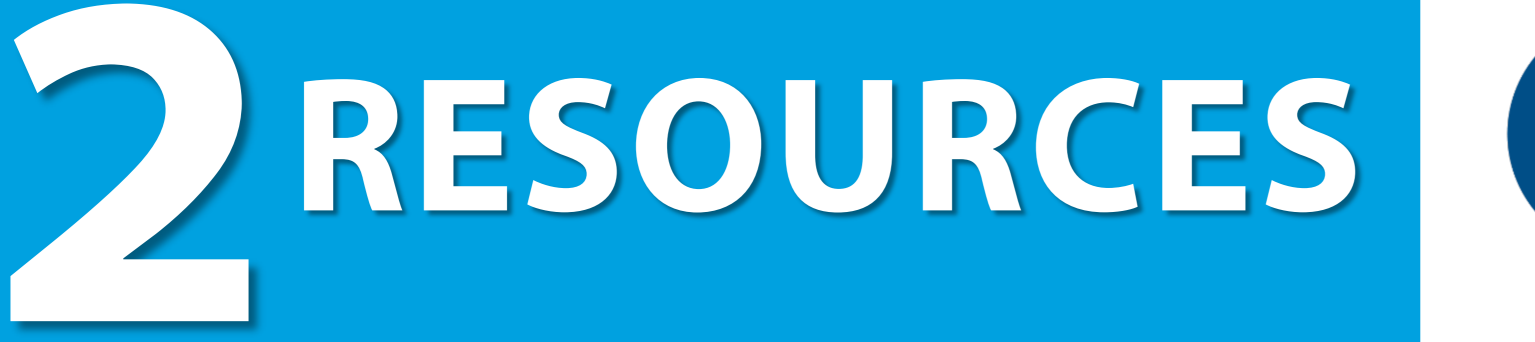
Within external contracts, the facility may carry out the following examinations: Computed Tomography, Magnetic Resonance Imaging, X-rays, PET, endoscopy, spirometry or other laboratory test.

The terms of the contracts determine the fast time to complete the procedures, which facilitates diagnosis and eventual AE checking at a rapid pace in patients from clinical trials.













CLINICAL TRIAL TEAM





10 RESEARCHES















CONSTANT RESEARCHES TEAM



prof. dr n. med. Andrzej Potemkowski

neurologist

22 years of experience in

clinical trials/

67 clinical trials



dr n. med. Aneta Wódecka

neurologist

8 years of experience in clinical trials/

20 clinical trials

dr n. med. Marcin Ratajczak

neurologist

- 16 years of experience in
 - clinical trials/
 - **78 clinical trials**



lek. Anna Ratajczak

pediatrician

16 years of experience in clinical trials/ **47 clinical trials**

lek. Adam Łopuszko

psychiatrist

6 months of experience in clinical trials/

6 clinical trials



COORDINATORS



mgr inż. Nina Omelańska

8 years of experience in clinical trials/

55 clinical trials



mgr Beata Miąsek 8 years of experience in clinical trials /



mgr inż. Agata Piłatowska

5 years of experience in clinical trials /

36 clinical trials



mgr Jakub Burdanowski

4 years of experience in clinical trials /

28 clinical trials

41 clinical trials



mgr Magdalena Drąg

6 years of experience in clinical trials /

22 clinical trials



mgr Maciej Kaczmarek

4 years of experience in clinical trials / **30 clinical trials**



dr n. med. Małgorzata Ryder-Woźniak

2 years of experience in clinical trials /

20 clinical trials



NURSES



Izabela Zamilska

12 years of experience in clinical trials /

72 clinical trials



mgr Kinga Bakalarczyk

2 years of experience in clinical trials / **17 clinical trials**



mgr Anna Kozińska

7 years of experience in clinical trials /

54 clinical trials



mgr Jolanta Rudzińska

6 years of experience in clinical trials /

38 clinical trials



4 years of experience in clinical trials /

28 clinical trials



mgr Paulina Ceglarz 2 years of experience in clinical trials / **3 clinical trials**



RATERS



dr n. med. Jerzy Kolenda

14 years of experience in clinical trials /

52 clinical trials



mgr Anna lwińska

2 years of experience in clinical trials /

7 clinical trials



mgr Malwina Prajwowska

4 years of experience in clinical trials /

11 clinical trials



mgr Agata Zapała

7 years of experience in clinical trials /

15 clinical trials



mgr Emilia Bastrzyk

6 monts of experience in clinical trials /

6 clinical trials



mgr Paulina Sobierajska

6 monts of experience in clinical trials /

6 clinical trials



mgr Paulina Rekowska

6 monts of experience in clinical trials /6 clinical trials





- SOP (STANDARD OPERATING PROCEDURES) A SET OF DETAILED WRITTEN INSTRUCTIONS, STEP BY STEP, TO ACHIEVE THE UNITY OF A SPECIFIC FUNCTION
- QUALITY CONTROL

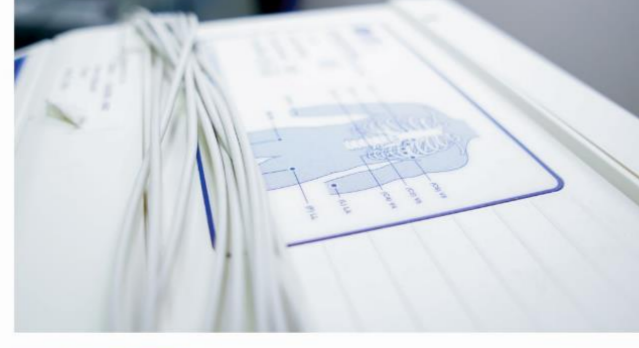




EQUIPMENT

We have certified equipment with a technical passports and full documentation.

- FREEZER UP TO (-80°C)
- FREEZERS UP TO (-25°C)
- LABORATORY CENTRIFUGES
- **COOLED CETRIFUGE UP TO DO 2°C**
- **REFRIGERATORS FROM 2°C TO 8°C**
- ECG
- DEFIBRILLATOR
- **SPHYGMOMANOMETERS**
- INFUSION PUMPS
- WEIGHT WITH STADIOMETER
- LAMINAR CHAMBER













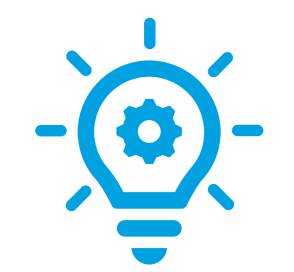




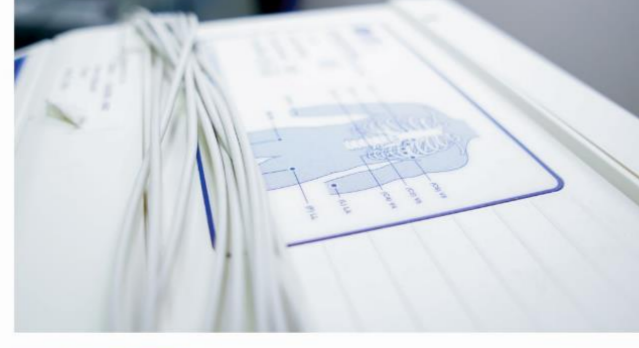




Our facility is also equipped with electronic system for temperature monitoring in medicines storehouse. The test product as well as biological samples are monitored.



The measurement is performed for 24 hours and notification about exceed is sent via text message or e-mail. The measurement is supervised by the rapid response team (up to 1 hour after the alarm occurred). It is possible to print reports on temperature control of any period of time.













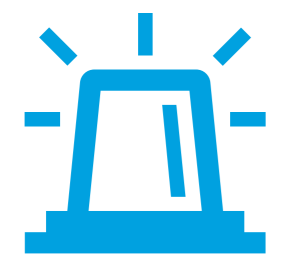


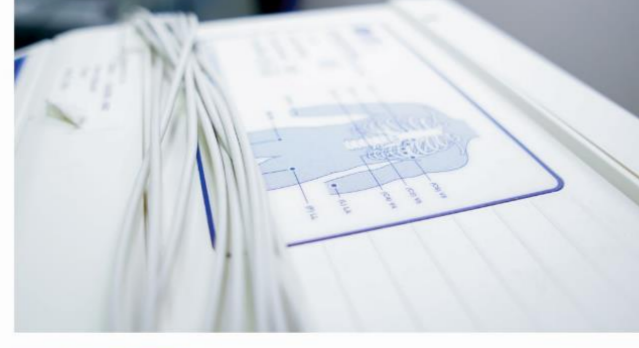






The facility has a power generator in the event of a power failure.



























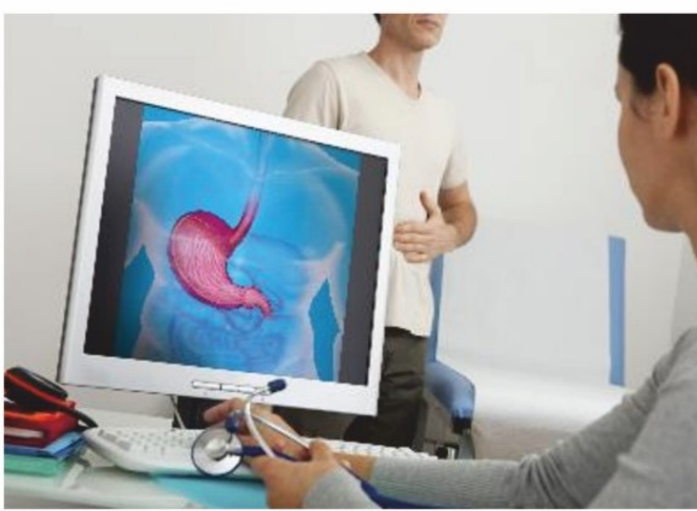
AREAS OF CLINICAL TRIALS

NEUROLOGY PSYCHIATRY DERMATOLOGY OPHTHALMOLOGY NEPHROLOGY GASTROENTEROLOGY DIABETOLOGY PRIMARY HEALTH CARE















COOPERATION

WE ARE OPEN TO NEW AREAS OF CLINICAL TRIALS!





102

CONDUCTED RESEARCH

9

TOP RECRUITER





1598

RECRUITED PATIENTS





1648

PATIENTS SUBJECTED TO SCREENING







COOPEARTING SPONSORS



COOPERATING CRO





16

AUDITS

FDA AUDIT







CURRENTLY CONDUCTED CLINICAL TRIALS





119

PATIENTS IN CURRENTLY CONDUCTED TRIALS



MONTHLY VISITS





OPERATING RULES







- TRAININGS

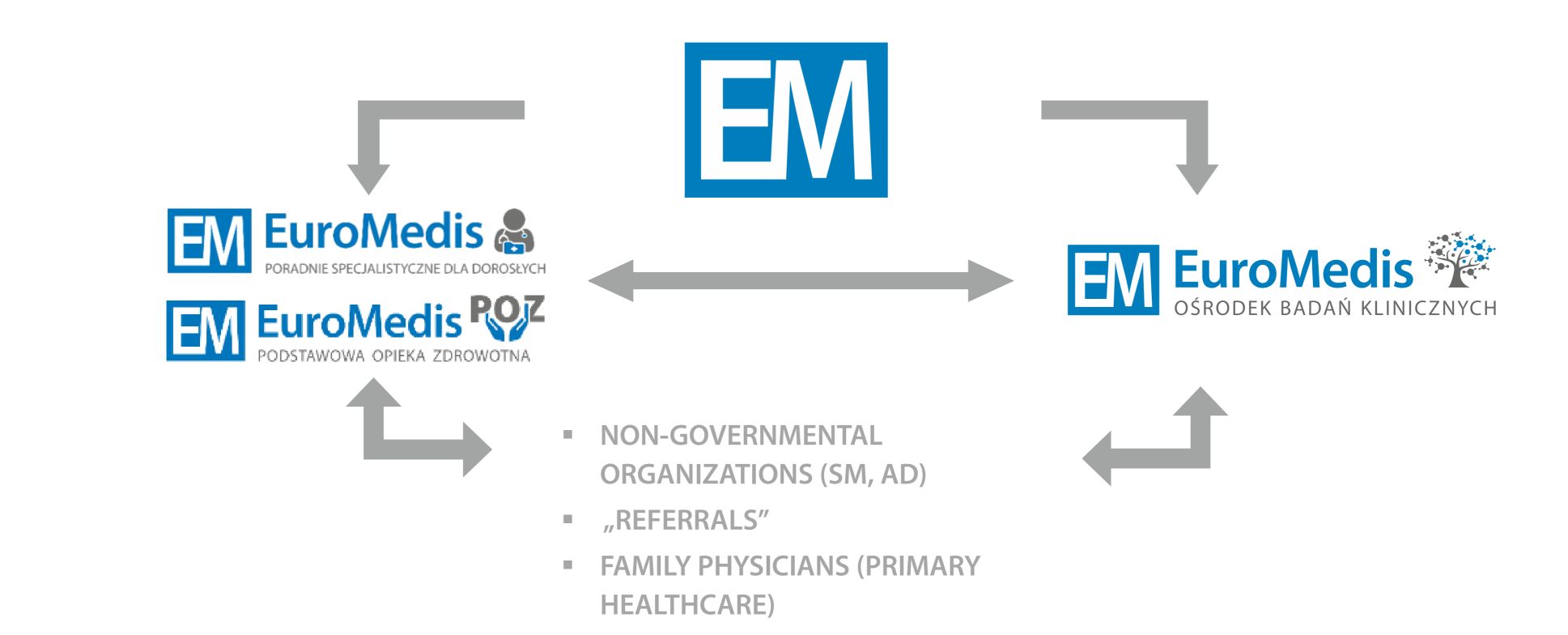
DISCUSSION OF EVERY INDIVIDUAL PATIENT'S CASE AND RELATING ISSUES OVERVIEW OF THE COURSE OF EACH CLINICAL TRIAL AND THE RELATED PROBLEMS

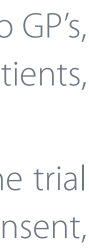




STRATEGIES IN RECRUITMENT

Connecting the medical activity provided by Specialistic Outpatient Clinics and Clinical Trial Centre in one facility enables constant flow of patients, it includes also GP's, 'referrals' and associations dealing with diseases on which they conduct clinical trials. It considerably increases recruitment possibilities from own database of patients, however, it also enables further care provided by the very same specialists once their participation in the trial has ended. All patients are informed about the possibility of participating in clinical trials as a treatment option. The patients who meet the criteria of being included in the trial and still are willing to take part, are invited to a special visit where all procedures and ICF – Informed Consent form are discussed. After the visit and patient's consent, we start the procedures connected with performing the given examination







- Region of north-west Poland: 3.000.000 people
- Szczecin and adjacent poviats: 600.000 people





STRATEGIES IN RECRUITMENT

EuroMedis runs a website devoted to clinical trials as well as numerous information campaigns in social media. We also prepare our own information materials regarding clinical trials.

EuroMedis is also an organizer of scientific seminars for patients and their guardians as well as scientific and information meetings for doctors.



information materials

www.EuroMedis.pl/ portal for doctors / social media

direct meetings/ conferences





FUTURE PLAN

- ELEKTRONIC MEDICAL DOCUMENTATION WITH REMOTE ACCESS TO SD VERIFICATION FOR CRA
- OWN MRI
- OWN WŁASNA ENDOSCOPY





WHY EUROMEDIS CLINICAL TRIAL CENTER?

- OVER 15 YEARS OF EXPERIENCE ON THE MARKET
- SPECIALIZED MEDICAL STAFF

²24/7 DIAGNOSTICS FOR PHASE 1 CLINICAL TRIALS

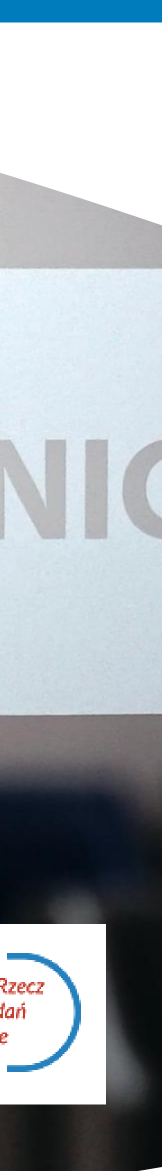
- EXPERIENCED CLINICAL TRIAL COORDINATORS
- DEDICATED TEAM FOR CONTACT WITH REFERRALS
- FULL DIAGNOSTIC AND LABORATORY FACILITIES
- 24/7 DIAGNOSTIC DEPARTMENT FOR 1 PHASE OF CLINICAL TRIALS
- MODERN TECHNOLOGICAL SOLUTIONS 24h
 TEMPERATURE MONITORING SYSTEM
- TELEPHONE CONTACT 24h AND COMPANY
 TRANSPORT FOR THE PATIENTS OF THE FACILITY
- QUALITY DEPARTMENT

BADANIA KLINI





Stowarzyszenie na Rzecz Dobrej Praktyki Badań Klinicznych w Polsce





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