CASE STUDY

Emergency ultrasound-assisted examination of skin and soft tissue infections in the paediatric emergency department.

CATEGORISING

<table>
<thead>
<tr>
<th>Type:</th>
<th>Clinical trial</th>
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</thead>
<tbody>
<tr>
<td>Subtype and Category:</td>
<td>Clinical trial with medical devices (according to ClinO-MD)</td>
</tr>
<tr>
<td>Category:</td>
<td>C1 not conformity-related</td>
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</tbody>
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BACKGROUND

This study evaluated the test characteristics of clinical examinations of paediatric patients, supplemented with bedside emergency ultrasound, and compared them to clinical examination alone, to identify skin and soft tissue infections that require drainage. The ultrasound device bears a CE mark, but is not used as intended. Since the bedside emergency ultrasound was added to clinical practice and the investigation of the clinical performance of adding bedside ultrasound was the goal of the study, the study qualifies as a clinical trial (and not as an observational study according to HRO chapter 2). - The study is conducted by an academic sponsor, who does not intend to use the data generated in the study to establish or demonstrate the conformity of the device for the new purpose investigated in the study.

METHODS

This was a prospective study of clinical examinations, supplemented with bedside emergency ultrasound, as a diagnostic test to evaluate patients 2 months to 19 years old. These patients were evaluated for skin and soft tissue infections in a paediatric emergency department. Each patient was clinically examined by the responsible physician (standard) and by an independent radiologist, who performed a bedside emergency ultrasound (result not communicated to the responsible physician). The reference standard that determined if a lesion required drainage was defined as “pus expressed at the time of the emergency department visit or within 2 days by follow-up assessment”. The primary outcome was the degree of agreement between clinical exam alone, and clinical exam supplemented with ultrasound, on the number of lesions that required drainage within 2 weeks after clinical examination. No other study-related procedures were performed except for the ultrasound.

QUESTIONS OF THE CATEGISER

**Does the research project come under the scope of application of the Human Research Act?**

Yes

**BECAUSE**

This project was based on a study protocol that defines the exact procedures to be used. It included a relatively large number of persons and was not based on individual cases (“method-driven search for generalizable knowledge”, defined as research by HRA). Children (“persons”) aged 2 months to 19 years, who suffered from skin and soft tissue infections (“research concerning human diseases”) were clinically examined and were given bedside emergency ultrasound.

**Is the research project a project involving living persons?**

Yes

**BECAUSE**

Children (“persons”) aged 2 months to 19 years, who suffered from skin and soft tissue infections, were included in this research project.
Is the research project a clinical trial?
Yes

BECAUSE
The performance of an ultrasound device ("medical device") was systematically tested in children ("persons") aged 2 months to 19 years, who suffered from skin and soft tissue lesions. The goal was to examine how consistent two examination techniques were in diagnosing lesions that require drainage.

Does the trial involve investigating a medical device (in vitro diagnostics excluded) or any other device as defined in Article 1 of the Medical Devices Ordinance of July 1, 2020?
Yes

BECAUSE
The performance of an ultrasound device ("medical device") was systematically tested in children ("persons").