Orthopedic registries have been kept for half a century (1,2). They have created a powerful model for evidence-based arthroplasty and surveillance. Spine surgery is widespread and the numbers are increasing. There is little research into the harm or benefit of spine surgery. The few available randomized trials have been criticized for poor generalizability (3). Registries are pragmatic, long-term observational studies with large numbers. This study aimed to describe the perceptions of decision-makers in major orthopedic centers regarding the value, implementation and use of spine surgery registries.

A 33-item survey was sent to CEOs and heads of spine surgery of the International Society of Orthopedic Centers (ISOC). ISOC includes 21 hospitals worldwide with a special focus on high-quality musculoskeletal care. Twelve out of 20 member centers (60%) replied to the survey. Seven have working registries; five in Europe and two in North America. The estimations for the cost/year were distributed more evenly: $10,000 [2], $20,000 [1], $50,000 [1]. Society cannot afford unnecessary surgery nor renounce to cure patients with effective treatments. Spine surgery registries provide high levels of evidence. The cost of implementing a registry is limited in comparison to RCTs. Spine registries can pragmatically fill our knowledge gap by turning every operated patient into a study participant.

Keywords: Spine; research; management; outcomes; prospective studies; registry; observational

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survey. Seven have working registries; 5 in Europe and 2 in North America. The oldest one started in Sweden in 1993 and the most recent in Italy in 2015 (Table 1). Allocated human resources (person-hours per week) varied from 0–5 in the Swedish registry, to more than 50 hours in the Italian registry. Most registries allocated 10 to 40 person-hours weekly. Five registries were combined electronic and paper; two were only electronic. Six were a combination of internal and external registries; 1 institution had an external registry (Table 2).

Five institutions reported not having a spinal registry. The reasons were funding (4), reduced case-load (2), organizational difficulties (3). Two institutions considered a spine registry a priority in the short-middle term, whereas 3 institutions considered it a priority for the long term. Finally, 2 institutions estimated the launching cost to be around $10,000, whereas 2 institutions estimated the cost to be $50,000. The estimations for the cost/year were distributed more evenly: $10,000 [2], $20,000 [1], $50,000 [1].

The perceived value of the spinal registry was high among both CEOs and Heads of Spine Surgery. Interestingly, clinicians perceived that their staff were less satisfied with the registry than they were, while the CEOs felt the staff was more satisfied than they were. This could reflect a more realistic perception (from clinicians) of the burden of work related to a registry. More experienced registries were less resource-consuming, though all registries consumed specific human resources. Limiting factors to implementation of registries were cost and organizational difficulties. The perceived cost of registries (reflecting the willingness to allocate resources) was modest (around $10,000 per year).

Surgery registries, and specifically spine registries, provide high levels of evidence in an area in which RCTs are often difficult to execute or unfeasible for ethical reasons (5). The cost of implementing a registry is limited in comparison to RCTs, whereas the information obtained from the registries is of high quality and generalizable (6). Understanding the specific advantages and disadvantages of keeping a spine registry and the cooperation with institutions that already have registries will help to further develop registries not only as valuable research tools but also as a means to achieve the highest standard of care.

Spinal disease is a burden. Society cannot afford unnecessary surgery nor renounce to cure patients with effective treatments. Spine registries can pragmatically fill our knowledge gap by turning every operated patient into a study participant.

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Footnote
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