

# Towards integrated pediatric care; solutions, effectiveness and safety considerations

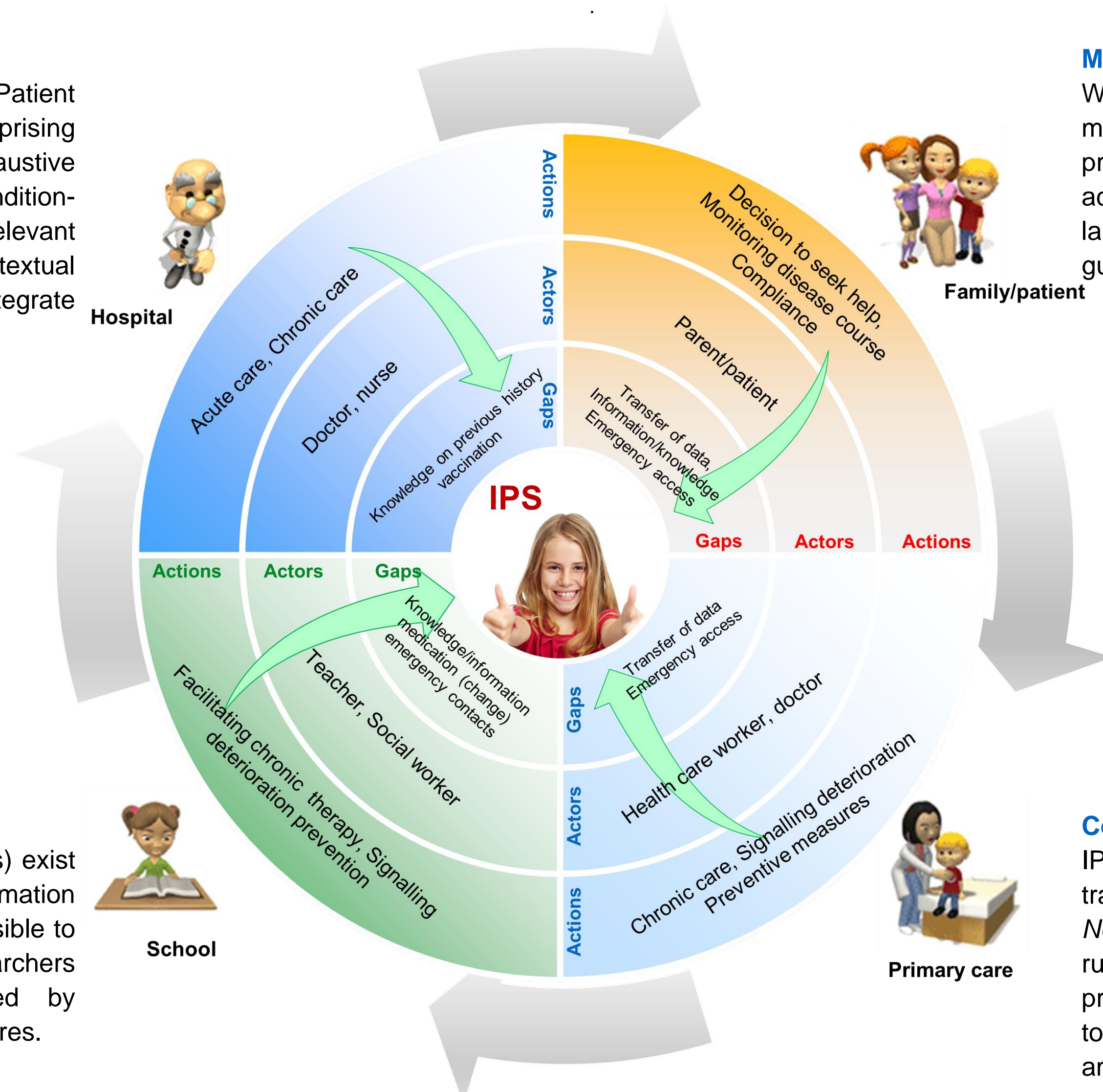
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## AIM

to study if the International Patient Summary (IPS) standard comprising a minimal and non-exhaustive specialty-agnostic, condition-independent, but clinically-relevant collection of clinical and contextual information, can be used to integrate health data for children

## Methods

We described pathways identifying main components or care covering preventive, family, supportive, and acute care. Data were retrieved from large collaborative studies and guidelines.



## Results

Electronic health record (EHRs) exist at focal points but lack information transfer. Resources are accessible to health policy makers or researchers mostly. Data are influenced by variability of health care structures.

## Conclusions

IPS could link health data at transition points for prompt feedback. Needs rules on what data to use and how to present them across the pathway to empower patients to access and amend IPS data



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