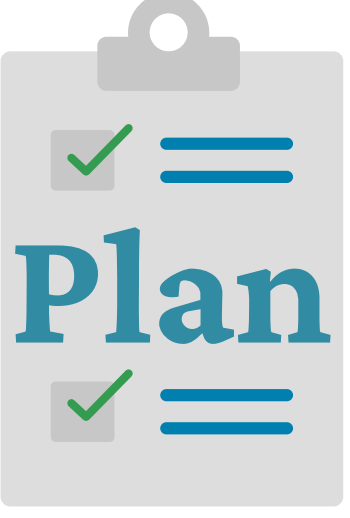


# Pregnancy in rCTDs

## 5 things you should know

**1**  **Plan** your pregnancy with your doctor

**2** **Discuss** treatment strategy with your doctor



**3** **Talk** to your doctor before stopping any medication



**4** **Inform** yourself with reliable sources



**5** You are not alone, **reach out!**



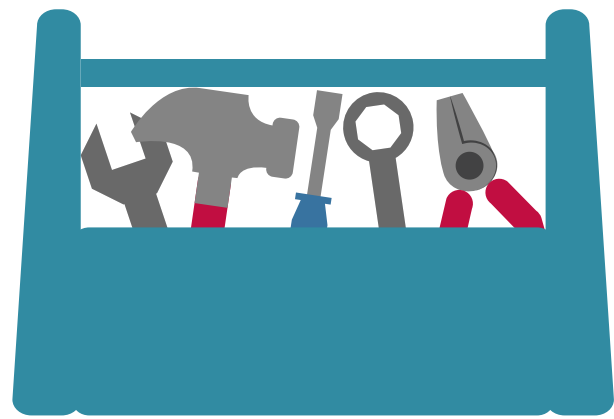
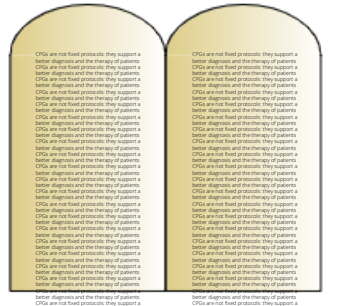
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- [ERN ReCONNET webinar on pregnancy](#)
  - blablabla
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# Clinical Practice Guidelines

## What you need to know



CPGs are not fixed protocols: they support a **better diagnosis** and the therapy of patients



CPGs in rare diseases are very important and when there isn't sufficient evidence **other tools** are useful



ERNs can enhance the **development** of CPGs



Patients can play an **active and crucial role** in the development of guidelines



**active and crucial role** of guidelines



**Lay versions** are essential for informed choices



- ERN ReCONNECT webinar on Clinical Practice Guidelines
- blablablaba
- State of the art on Clinical Practice Guidelines
- blabla
- bla

This infographic was made by the ERN ReCONNECT ePAGs. You can print it out, download it, disseminate it however you want.

[ERN ReCONNECT](#) is the European Reference Network on Rare and Complex Connective Tissue and Musculoskeletal Diseases. [ePAGs](#) are the European Patient Advocacy Groups that bring together elected patient advocates and affiliated organisations in order to ensure that the patient voice is heard throughout the ERN development process.