ELF- Levels of involvement in CRCs

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| Level 1: Low involvement: no charge |
| * Advice on how to involve patients in the CRC * Introducing CRCs to relevant patient organisations and advising on patient involvement * Introducing CRC to someone already in our network * Basic information about the CRC cascaded to our network * Providing templates for patient information * Basic information about the CRC’s activity and results on the ELF website and social media * Review of final publications |
| Level 2: Medium involvement: approx. €5000 per year |
| As for Level 1 but also including:   * Participation in relevant CRC meetings and groups * Formation of and recruitment to a Patient Advisory Group and ongoing co-ordination of and consultation with the Patient Advisory Group, including training and support, and linking with CRC. This can include basic information gathering from the group, such as their views on patient priorities, study design and data collection. * Integration of patient perspectives into publications * Lay summaries of the CRC activity, outcome and publications on the ELF website and social media * A factsheet or lay version of the CRCs output |
| Level 3: High involvement: additional cost per activity |
| As for levels 1 and 2 but also including:   * Patient-centred literature review – approx. €2000 * Creation and dissemination of patient surveys and analysis of results – approx. €4000 per survey, depending on size and reach (excluding translation costs) * Focus groups with national patient organisations – approx. €1000 per group depending on size * Co-produced patient information materials – approx. €1000 per document, depending on size (excluding translation costs) |