

CORRIGENDA

Genetic epidemiology, prevalence, and genotype–phenotype correlations in the Swedish population with osteogenesis imperfecta

Katarina Lindahl, Eva Åström, Carl-Johan Rubin, Giedre Grigelioniene, Barbro Malmgren, Östen Ljunggren and Andreas Kindmark

European Journal of Human Genetics (2015) **23**, 1112; doi:10.1038/ejhg.2015129

Correction to: *European Journal of Human Genetics* (2015) **23**, 1042–1050; doi: 10.1038/ejhg.2015.81; published online 6 May 2015
 Post online publication, the authors have chosen to add some information to Supplementary Tables 1 and 2. The original files have been replaced with their updated counterparts and are available to view online.

A prospective cohort study assessing clinical referral management & workforce allocation within a UK regional medical genetics service

Caroline Benjamin, Catherine Houghton, Claire Foo, Chris Edgar, Gail Mannion, Jan Birch, Ian Ellis and Astrid Weber

European Journal of Human Genetics (2015) **23**, 1112; doi:10.1038/ejhg.201582

Correction to: *European Journal of Human Genetics* (2015) **23**, 996–1003; doi:10.1038/ejhg.2015.33; published online 11 March 2015
 Corrections have been made to Figure 1 and a revised copy of this paper appears in this issue.

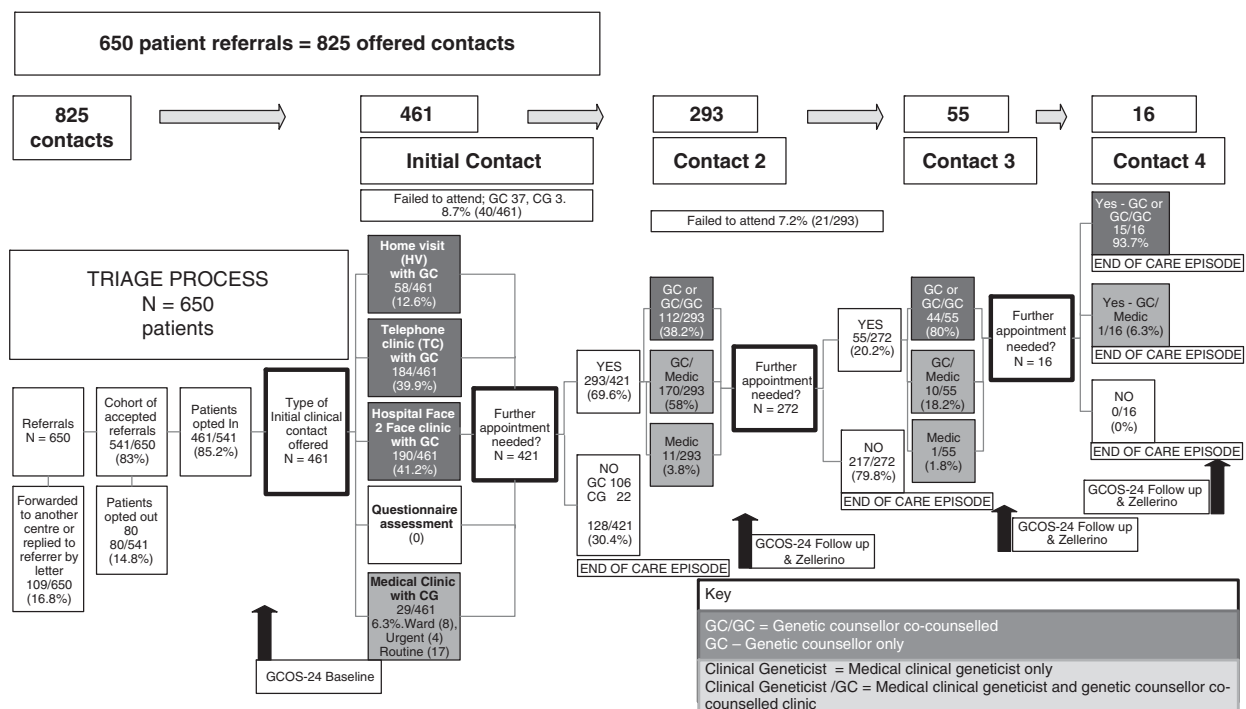


Figure 1 Progress of the 650 new patients referred over the 3-month period (12.12.2011–12.03.2012), resulting in 825 offered contacts – with 12 months of follow-up until 12.03.2013.