

primary studies - published RCT

Improved treatment response to dornase alfa in cystic fibrosis patients using controlled inhalation.

Code: PM21737560 Year: 2011 Date: 2011 Author: Bakker EM

Study design (if review, criteria of inclusion for studies)

multicentre, double-blind, randomised controlled clinical trial

Participants

cystic fibrosis patients on maintenance treatment with 2.5 mL dornase alfa once daily

Interventions

patients on maintenance treatment with 2.5 mL dornase alfa once daily were switched to a smart nebuliser and randomised to small airway deposition (n = 24) or large airway deposition (n = 25) for 4 weeks.

Outcome measures

The primary outcome parameter was forced expiratory flow at 75% of forced vital capacity (FEF(75%)).

Main results

FEF(75%) increased significantly by 0.7 sd (5.2% predicted) in the large airways group and 1.2 sd (8.8% pred) in the small airways group. Intention-to-treat analysis did not show a significant difference in treatment effect between groups. Per-protocol analysis, excluding patients not completing the trial or with adherence

Authors' conclusions

Improved delivery of dornase alfa using a smart nebuliser that aids patients in correct inhalation technique resulted in significant improvement of FEF(75%) in children with stable cystic fibrosis. Adherent children showed a larger treatment response for small airway deposition.

http://dx.doi.org/10.1183/09031936.00006211

See also

Eur Respir J. 2011 Dec;38(6):1328-35. Epub 2011 Jul 7.

Keywords

Deoxyribonuclease; Airway clearance drugs -expectorants- mucolytic- mucociliary-; pharmacological_intervention; Respiratory System Agents; non pharmacological intervention - devices OR physiotherapy; Inhalation OR nebulised; nebuliser; Dornase alpha; Pulmozyme;