Starting HMV at home: a reasonable option for many patients?

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Abstract

Background and objective In the current study, we undertook a more detailed exploration of the reasons why patients undergoing HMV were screened out of a recently published study in order to better understand how applicable home initiation of HMV is under real life conditions.

Methods All referred patients who had an indication for starting HMV were screened to participate in the Homerun study. In this trial 512 patients were screened out of the study. Those patients not enrolled in the trial were divided into the following 3 groups: (1) those not fulfilling the inclusion criteria; (2) those meeting the exclusion criteria and 3) those excluded on the basis of medical or organisation reasons. Each group was then further divided into those who would likely have been suitable for initiation of HMV at home in real world practice and those who were unsuitable.

Results Based on inclusion criteria (group 1) 116 patients could not start HMV in real life, while this was 245 patients in the study. Based on the exclusion criteria (group 2) 11 patients could not start in real life while this was 79 in the study. One hundred and eighty-eight could not be enrolled in the study due to medical and organisational reasons (group 3), while in real life this was only 95.

Conclusion This study indicates that more than 55% of patients who did not participate in the Homerun study could have started HMV at home in real life.

Keywords Home mechanical ventilation, Telemonitoring, Patient selection, Health care organization
Introduction
Home mechanical ventilation (HMV) is an effective treatment for patients with chronic hypercapnic respiratory failure, resulting in improved clinical symptoms, health status and survival in patients with both restrictive and obstructive pulmonary disorders [1]. Consequently, we currently see that rapid growth in the number of patients on HMV is occurring [1]. In the Netherlands, it is common to initiate HMV in an inpatient setting with monitoring of gas exchange; however, the observed increase in demand for HMV represents an extra burden to the hospitals with regard to the availability of hospital beds. In addition, hospitalisation is stressful for patients as they have to leave their social environment.

More importantly, the existing patient tailored care system at home in every individual patient, often a combined team of professional and informal caregivers, will not be positively affected by hospitalisation. Therefore, there is a need for alternative ways of initiating HMV.

With this in mind, we designed a national study, The Dutch Homerun trial, [2] based on previously designed local studies on home initiation of HMV, [3, 4] in which we compared initiation of HMV at home with inpatient initiation. We showed that home initiation was equally effective as hospital initiation. However, a large group of patients did not meet the inclusion criteria (84%), therefore, they did not get the opportunity to start ventilatory support at home.

Therefore it is important to know the number of patients that could be candidate for initiate HMV at home. So we need to know if subjects were solely excluded due the study design and strict enrolment criteria or as a consequence of truly being unsuitable for home initiation of NIV based on clinical or social factors. The findings could have a significant impact on HMV protocols for initiating HMV in the home setting with implications for resource and personnel allocation in the future. Therefore, we undertook this study to identify and review the reasons why patients were excluded from the Homerun study to better understand the number of individuals in usual clinical practice who may be eligible for initiation of NIV in the home setting.

Methods
During the inclusion period of the Homerun study, all patients who qualified for HMV were screened. For participation, they had to meet the following inclusion criteria: diagnosis of neuromuscular disease (NMD) or thoracic cage disorder with the indication for HMV based on complaints of alveolar hypoventilation (fatigue, headache of dyspnoea) combined with all following criteria: [2].

- Arterial carbon dioxide > 6.0 kPa daytime, or arterial or transcutaneous carbon dioxide > 6.0 kPa at night, or orthopnea as a result of diaphragm paralysis.
- Over 18 years of age.
- In the opinion of the supervising centre for HMV, the patient had a sufficient social or professional network to permit the safe initiation of HMV in the home.

The following exclusion criteria were applied:
- Previously received CPAP therapy or non-invasive ventilatory support.
- Requiring invasive ventilatory support.
- Living in a nursing home.

We categorised all non-eligible participants into 3 main groups by the following order: (i) those who did not meet the inclusion criteria (ii) those who met an exclusion criterion and (iii) those excluded from the study based on comorbidities or organisational reasons. The main groups were further subcategorized (see Fig. 1). The reason for not participating was evaluated to determine if the exclusion was solely based on the Homerun study design or whether there was a decisive factor that prevented the save initiation of HMV at home.

Each of these 3 groups were then further divided into those in whom it was felt exclusion from the study was solely based on the Homerun study design, and those in whom there was another reason for preclusion from the ability to initiate HMV. We use the term real life for the patient situation out of the study designed factors for not participating and into barriers which prevent from initiation at home in a daily practice work situation.

Results
During the recruitment period of the Homerun study, a total of 608 patients were screened and commenced on HMV. However, only 96 participants were eligible for the study, leaving 512 patients to be set up on HMV outside of the study. The reasons for not participating were categorized into three main groups (as mentioned in the Methods section). These groups were further subdivided to gain detailed insight (see Fig. 1).

Patients not fulfilling the inclusion criteria
Two-hundred and forty-five participants, making up 48% of excluded participants, did not fulfil the inclusion criteria, primarily due to not having a NMD or TCA. However, other disorders characterised by hypoventilation might still be considered suitable candidates to start HMV at home in real life. Age limitation, an insufficient supportive network, and lack of motivation would remain reasons for not initiating HMV at home in real life (Fig. 2). Especially in ALS patient, although there is a normocapnia and absence of orthopnea there can be reasons, as experienced extensive work of breathing...
or frequent pulmonary infections, to start HMV in this exceptional cases.

**Patients fulfilling the exclusion criteria**
In the study 15% met the exclusion criteria, seventy-nine participants. From the 3 mentioned exclusion criteria, only the need for invasive ventilation renders starting HMV at home would be impossible in real life (Fig. 3).

**Comorbidity and organisational reasons for not participating**
In this group a total of one hundred and eighty-eight patients had reasons for not participating. Due to logistical issues, 85 patients started HMV under guidance and supervision of an academic center of home mechanical ventilation in-hospital but outside one of the 4 HMV hospitals and could therefore not be randomized. Of note, if these patients met the inclusion criteria and would not have been excluded based on exclusion criteria, they could have initiated HMV at home. Unfortunately, inclusion in the Homerun trial was not discussed with these patients. The other reasons for non-participation related to comorbidity and organisational issues are presented in Fig. 4.

In summary, while 512 patients were excluded from the Homerun study, in real life up to 290 of these could have started HMV at home, with the remaining 222 unsuitable of initiation of therapy at home due to various reasons (Fig. 5).
Discussion

This study shows that initiation of HMV at home can be offered to a larger group of patients than was initially thought based on the Homerun study results. This is important as the Homerun study suggests that starting HMV at home is only feasible in a very select group of patients with an NMD or thoracic cage problem, this suggestion could be extended to the study of Hazenberg which used the same inclusion and exclusion criteria [2, 3].

Based on a recent study by Duiverman and colleagues, it has been shown chronic obstructive pulmonary disease (COPD) patients, which represent 13% of patients being referred for HMV, can be commenced on HMV at home as well [4]. Considering the other “Inclusion-related factors” mentioned in group 1, that might prohibit starting HMV at home, it remains difficult to upgrade an insufficient social network or improve patient motivation. While staying at home for starting therapy could positively influence the motivation for therapy, although we have no proof for this statement. Other actions to make the initiation at home more attractive with regards to motivation could be obtained by education and raising more awareness that starting HMV at home is safe and equally effective as compared to an inpatient start.

Fig. 2 Distribution of patients fulfilling the inclusion criteria of the Homerun study versus real life

To justify HMV initiation in patients under 18 years of age, similar studies as the Homerun trial need to be carried out in younger patients. Lastly, the exact reasons why patients declined to participate in the study were not obtained. Better communication of the positive results combined with sharing the experiences of other patients might take away the possible burden(s) anticipated by these patients.

With regard to the exclusion criteria, it is obvious that the start of tracheostomized ventilation and its complexity of care is only feasible in a high care unit. However, other criteria for exclusion from the Homerun study do not make initiation of HMV at home impossible in real life. A Starting HMV in a nursing home is also feasible as nurses are available 24/7 and care can be provided instantly.

The last category for not participating is more complex and diverse as shown in Fig. 4, but two reasons are relevant for discussion. During the Homerun trial, 85 patients started HMV in a regional hospital, under guidance and supervision of an academic center of home mechanical ventilation and inclusion in the Homerun study was not discussed as a possibility; it is highly conceivable that at least some of these patients would have
qualified for HMV initiation at home. Regarding patients exclusion due to acute initiation of NIV, we know some patients may deteriorate acutely due to an infection, while other patients may gradually worsen over weeks or months. This latter group may have benefitted from home-initiated ventilation if identified and referred to a HMV centre at an earlier stage of their disease process. Education and improved awareness of health care professionals regarding positive effects of HMV in patients with or at risk of hypoventilation due to NMD or TD is important to reduce the number of patients needing start HMV in an acute setting.

Comparing this current study with previous work is difficult as we found only one study comparing hospital adaptation to HMV versus home adaptation [5]. While this observational prospective study showed improvements in both groups in terms of arterial blood gases and quality of life, however, it did not investigate reasons behind non-participation [5]. Several studies have investigated initiation of HMV in an outpatient setting compared to inpatient HMV initiation. Although reasons for not initiating HMV as an outpatient service were identified, it was unclear whether these reasons were primarily study-related or due to real-life circumstances. [6–9].

**Conclusion**

This study suggests that up to 55% of patients who did not participate in the Homerun study could have started HMV at home in real life. Based on the expected growth of patients who need HMV in the future, we should reconsider our current clinical HMV initiation process and start HMV at home rather than at the hospital whenever a patient is considered eligible based on real-life criteria. Investments in the telemonitoring, proper training of staff, and organisational restructuring are needed to make this happen.

**Fig. 3** Distribution of patients meeting the exclusion criteria for the Homerun study versus real life
Fig. 4 Comorbidity and organisational reasons for not participating in the Homerun study vs. real life

Fig. 5 Summary of reasons for not starting HMV at home by category for the Homerun study vs. real life

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Author contributions
R.B. and P.W were the primary authors of the paper. A.H., N.C. and M.G. assisted with data collection, which was confirmed by R.B. All authors were involved in interpretation of data and contributed to critical revision of important intellectual content and final approval of the version to be published.

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**Availability of data and material**
The datasets used and analyzed during the current study are available from the corresponding author on reasonable request.

**Declarations**

**Statements of ethics**
The data described in this article is distracted from the original study data which is already published. This original study was approved by the UMCG institutional review board (approval number 2014/529). Monitoring was performed on all sites by an external monitor, and a Data Safety Monitoring Board was installed. Individuals have given their written informed consent. Patients who were excluded from the original study no informed consent was obtained, this wasn't necessary, because only the reason for not participating was noted. All procedures were performed in accordance with relevant guidelines. The study was registered in the Dutch trial register (NTR4683) and later internationally (ClinicalTrials.Gov ID: NCT03203577).

**Consent for publication**
Patients included in the study were informed by the investigator that results would be published. The group of non-participants was logged but didn't consent for publication, because their specific situation for not participating in the study is categorized and therefore not seen as direct patient information and not involving or harmful for these patients. We see this data as management information for improving health care organization.

**Competing interests Statement**
Dr. van den Biggelaar reports personal fees from Philips, and Westfalen Medical B.V., both outside the submitted work. Dr. Hazenberg, Dr. Cobben, Dr. Gaytant and Dr. Gommers have nothing to disclose. Prof. Dr. Wijkstra reports grants from ZONMW; grants from VIVISOL, during the conduct of the study; grants and personal fees from Philips; grants and personal fees from RESMED, grants from Goedgegebuure, grants from vital air, personal fees from Bresotec, and personal fees from Synapse, outside the submitted work.

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