


# BMJ Open Quality Reducing low-value care: what can we learn from eight de-implementation studies in the Netherlands?

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**To cite:** Verkerk EW, van Dulmen SA, Westert GP, *et al.* Reducing low-value care: what can we learn from eight de-implementation studies in the Netherlands? *BMJ Open Quality* 2022;**11**:e001710. doi:10.1136/bmj-2021-001710

► Additional supplemental material is published online only. To view, please visit the journal online (<http://dx.doi.org/10.1136/bmj-2021-001710>).

Received 19 October 2021  
Accepted 23 August 2022



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## ABSTRACT

**Background** Reducing the overuse of care that is proven to be of low value increases the quality and safety of care. We aimed to identify lessons for reducing low-value care by looking at: (1) The effects of eight de-implementation projects. (2) The barriers and facilitators that emerged. (3) The experiences with the different components of the projects.

**Methods** We performed a process evaluation of eight multicentre projects aimed at reducing low-value care. We reported the quantitative outcomes of the eight projects on the volume of low-value care and performed a qualitative analysis of the project teams' experiences and evaluations. A total of 40 hospitals and 198 general practitioners participated.

**Results** Five out of eight projects resulted in a reduction of low-value care, ranging from 11.4% to 61.3%. The remaining three projects showed no effect. Six projects monitored balancing measures and observed no negative consequences of their strategy. The most important barriers were a lack of time, an inability to reassure the patient, a desire to meet the patient's wishes, financial considerations and a discomfort with uncertainty. The most important facilitators were support among clinicians, knowledge of the harms of low-value care and a growing consciousness that more is not always better. Repeated education and feedback for clinicians, patient information material and organisational changes were valued components of the strategy.

**Conclusions** Successfully reducing low-value care is possible in spite of the powerful barriers that oppose it. The projects managed to recruit many hospitals and general practices, with five of them achieving significant results without measuring negative consequences. Based on our findings, we offer practical recommendations for successfully reducing low-value care.

## INTRODUCTION

Reducing care that is proven to be of low value is a universal and persistent challenge.<sup>1</sup> Such low-value care, also called medical overuse, provides no, or very little, benefit to the patient if one takes into consideration its potential harm, costs, alternatives or patient preferences. In addition it also wastes resources.<sup>2</sup> The term de-implementation is increasingly being used to describe a move away from ineffective or harmful medical

## WHAT IS ALREADY KNOWN ON THIS TOPIC

- ⇒ Reducing low-value care can increase the quality and safety of care while maintaining or decreasing healthcare costs.
- ⇒ Reducing such care has proven to be difficult and knowledge about its de-implementation is scarce.

## WHAT THIS STUDY ADDS

- ⇒ This study shows that clinicians can reduce low-value care successfully, despite barriers such as a lack of time, financial considerations and the need to reassure patients.
- ⇒ We provide practical recommendations for de-implementation studies, such as only reduce low-value care that is supported by sufficient evidence, tailor a de-implementation strategy to counter the barriers, use repeated education and feedback for clinicians, and provide carefully developed patient information.

## HOW THIS STUDY MIGHT AFFECT RESEARCH, PRACTICE OR POLICY

- ⇒ More clinicians and researchers should reduce the overuse of low-value care, and our study will help them to improve their approach.

practices.<sup>3</sup> Such reduction of low-value care can increase the quality and safety of care. Hence, many initiatives have started worldwide, such as the Choosing Wisely campaign that began in the USA in 2012 and since then has spread to over 20 countries.<sup>4</sup> However, reducing low-value care has proven to be difficult and knowledge about de-implementation is scarce.<sup>5</sup> Two evaluations of Choosing Wisely recommendations showed marginal and varying results 1.5 years and 2.5 years after their release.<sup>6,7</sup> Literature reviews suggest that strategies comprising different components, addressing patients and clinicians, have the potential to reduce overuse.<sup>2,8</sup> However, the underlying mechanism in play is unclear and further experimentation and evaluation is needed.<sup>2,9</sup>

Several publications describe lessons learnt so far from de-implementation. A review



stated that involving physicians from the beginning is of great importance.<sup>10</sup> Another study evaluated eight de-implementation projects in a hospital and found that support from the hospital board was a key to their success.<sup>11</sup> An interview study among Choosing Wisely team members found that harm reduction is a significant motivator to reduce low-value care and that data collection could be challenging.<sup>12</sup> Further in-depth knowledge and experience of de-implementation, including its impact and the barriers and facilitating factors involved, is needed to determine what is necessary for successful de-implementation.<sup>13</sup>

In 2015, we started a nationwide programme in the Netherlands, comprising eight multicentre de-implementation projects that we prospectively monitored and evaluated. Each de-implementation project aimed to reduce a different type of low-value care. The projects were led by clinicians and set in multiple hospitals or primary care practices. This paper describes the lessons learnt from these projects and aims to contribute to the knowledge on de-implementation in clinical practice by answering three questions:

- ▶ What effects can be achieved by a multicentre de-implementation project?
- ▶ What barriers and facilitating factors might be encountered in de-implementation?
- ▶ What are the effective components of a de-implementation project, and why?

## METHODS

We prospectively monitored and evaluated eight multicentre de-implementation projects in the Netherlands from June 2016 to October 2018. This study was part of a national programme called 'To do or not to do? Reducing low-value care', described in [box 1](#) and in more detail in online supplemental file 1.

### The projects' structure

An overview of the projects can be found in [table 1](#). Six projects aimed at reducing low-value hospital care and two projects focused on low-value primary care. Each project leader chose a design and approach that would fit their project best, resulting in a diversity of study designs and

strategies. All projects evaluated the effect of the de-implementation strategy on the delivery of care. Six projects also measured the unintended effects of the strategy on patient outcomes and/or the use of other care. All project teams performed a structured process evaluation, and all projects obtained ethical approval before the start of their study. Several projects are described in more detail in other papers.<sup>14–23</sup>

### Evaluation

We used the Medical Research Council framework for process evaluation of complex interventions.<sup>24</sup> This framework helps to analyse why, and how, the planned intervention has led to the effect observed. Using this framework, we evaluated three components of the projects: (1) The effects of the projects on clinical practice; (2) The contextual barriers and facilitating factors that emerged; and (3) The experiences of the project leaders and the participating clinicians and patients with the different components of the projects.

For the first component, we report the quantitative outcomes of the eight projects on the volume of low-value care and on other outcomes that were measured. For components 2 and 3, EWV performed a qualitative analysis using Atlas.ti V.8.4.20 of the project teams' experiences and evaluations. We collected data on this using logbooks, reports and interviews. The project teams kept a logbook and delivered a report on their results and evaluation, for which they used a variety of qualitative and quantitative methods ([table 1](#)). In May 2018, two researchers from the coordinating team (EWV and PH) interviewed the project leaders of the eight teams. Details on these interviews are reported in the Consolidated criteria for REporting Qualitative research checklist (online supplemental file 2). The interviews included open-ended questions about the barriers and facilitating factors, the project leaders' experiences with different components of their project, the lessons they have learnt and their advice for other project leaders. Reports of the audiotaped interviews were sent to the project leaders for correction and confirmation.

EWV analysed the information reported in the logbooks, reports and interviews. Barriers and facilitators were classified using the framework of the determinants of change.<sup>25</sup> This framework identifies individual health professional factors, patient factors, professional interactions, incentives and resources, and social, political and legal factors. These categories were used for coding. We added one category (low-value care related) and three subcategories (interaction with patient, interaction with clinician, patient environment) to this framework. This was because some factors that we identified did not fit in the existing categories. The coding and description of results were verified by PH and discussed until consensus was reached.

### Patient and public involvement

We analysed eight de-implementation projects, each of which involved patients in their problem analysis, process

#### Box 1 Programme characteristics

- ⇒ 'To do or not to do? Reducing low-value care' was a national programme, coordinated by the eight university hospitals in the Netherlands.
- ⇒ The programme was both top-down and bottom-up, supported by stakeholders and initiated and led by clinicians.
- ⇒ Eight de-implementation projects were selected from 42 proposals by an independent committee.
- ⇒ The eight de-implementation projects received support from a central team, comprising the authors of this paper. The projects all followed similar steps according to the GroL and Wensing Implementation of Change Model.<sup>25</sup>
- ⇒ The projects ran from 2016 to 2018.

**Table 1** Overview of the eight projects

Project	Reduction in the inappropriate use of:	Setting	Problem analysis data source	Design	De-implementation strategy	Effect evaluation data source	Process evaluation data source
1	Inhaled corticosteroids for patients with mild chronic obstructive pulmonary disease	Five primary care cooperation groups, with a total of five pharmacists and 40 general practices	Focus group interviews with clinicians and patients	A before-and-after study with a national control group	Education of GPs and pharmacists. Publications in patient and professional magazines. Selection of patients whose use of inhaled corticosteroids was potentially unnecessary. Patient information	National database	Survey among clinicians and patients
2	Surveillance CT scans for patients cured of lymphoma	Nine hospitals' haematology wards	A survey among clinicians and patients	A before-and-after study with a national control group	Education of haematologists. Patient information (leaflet). Presentation at a patient association conference	National database	Survey among clinicians
3	Knee arthroscopies and MRIs for orthopaedic patients aged 50 years or older	Thirteen orthopaedic centres	Interviews and surveys among clinicians and patients	A difference-in-difference design with a national control group	Appointing clinical champions. Education of orthopaedic specialists. Patient information (leaflet). Feedback	National database	Survey among clinicians
4	Intravenous and urinary catheters	Seven hospitals' internal medicine and non-surgical subspecialty wards	A survey among patients and observations in clinical practice	A before-after study with an interrupted time series analysis	Appointing clinical champions. Education of physicians and nurses. Use of educational materials (poster, pocket card). Patient information (leaflets). Competitive feedback. Changes in the structure of medical records	Patients' medical records	Observations in clinical practice
5	Vitamins D and B <sub>12</sub> tests	Twenty-six primary care health centres, with a total of 158 general practitioners	Experience from an earlier pilot study	Cluster randomised study comparing two interventions	Education of GPs and feedback in intervention groups A and B. Patient information (leaflet, video clip and poster) in intervention group B only	Regional database	Interviews with clinicians and patients
6	Diagnostic laboratory tests	Four hospitals' internal medicine wards	Experience from an earlier pilot study and a survey among clinicians	A before-after study with an interrupted time series analysis and a control group of 19 hospitals	Conferences for physicians. Increased supervision of residents. Education of physicians. Feedback. Changes in the ordering system	Hospital registries	Survey among clinicians

Continued

**Table 1** Continued

Project	Reduction in the inappropriate use of:	Setting	Problem analysis data source	Design	De-implementation strategy	Effect evaluation data source	Process evaluation data source
7	Surveillance visits for patients cured for basal cell carcinoma	Three hospitals' dermatology wards	Interviews and focus group interviews with clinicians and patients	An uncontrolled before-and-after study	Personalised patient information	A survey among patients	Interviews with clinicians and patients
8	Upper gastrointestinal endoscopies for dyspeptic patients	Four hospitals' gastroenterology wards	Focus group interviews with clinicians and patients	A randomised controlled trial	Interactive e-learning for patients	Patients' medical records	A survey among patients

evaluation or both (specified in [table 1](#)). Patients who had been involved in the problem analysis contributed to the development of the de-implementation strategy. In addition, a representative of the Dutch patient federation became a member of the programmes advisory board. This board regularly met and advised the coordinating team on the design and progress of the programme.

## RESULTS

First, we report the quantitative outcomes of the eight projects on the volume of care. Then, we report the results of our qualitative evaluation of the barriers and facilitating factors for de-implementation, and the experiences of the project leaders and the participating clinicians and patients with the different components of the projects.

### Effects on clinical practice

The quantitative effects of the projects are shown in online supplemental file 3 and summarised in the text below. Five projects (4, 5, 6, 7, 8) showed a positive effect of the de-implementation strategy, the reduction in low-value care ranging from 11.4% to 61.3%. Project 5 compared two interventions and found a larger reduction in the group that received the additional patient information (10% extra reduction for vitamin D and a non-significant extra reduction of 4% for vitamin B<sub>12</sub>). Project 6 and 8 also collected data from a concurrent control group and both found a larger reduction (reduction of 11.4% in project 6 and 61.3% in project 8) in the intervention group compared with the control group (increase of 2.4% in project 6 and reduction of 17.5% in project 8). The remaining projects 4 and 7 studied one intervention arm and no control group.

Three projects (1, 2, 3) found no effect of the de-implementation strategy. Project 1 found a significant reduction in the control group, compared with no difference in the intervention group. Project 2 found no change in both groups. Lastly, project 3 found a reduction in low-value care in both the intervention and the control groups, but no difference between these groups. Six projects monitored balancing measures and found no negative effects of the de-implementation on use of other care and patient outcomes.

### Barriers and facilitating factors for de-implementation

The project teams found multiple factors that either hindered or facilitated the de-implementation of their low-value care practices. All the factors are presented in online supplemental file 4. Below, we describe the most frequently reported factors.

#### Factors related to low-value care

Regarding the factors that relate to the low-value care, evidence and a consensus among clinicians were the most frequently mentioned factors. These factors both facilitated de-implementation when they were present, and hindered de-implementation when they were absent.

### Individual health professional factors

A major barrier related to individual health professionals was a lack of knowledge about the low-value care. The knowledge and a belief that the care's harms outweighed its benefits facilitated de-implementation. For example, receiving a reminder of the fact that urinary catheters cause discomfort and lead to infections motivated clinicians to remove them more promptly. Another major barrier is the clinicians' fear of missing disease, and discomfort with uncertainty. In addition, clinicians felt that by providing low-value care they were meeting their patient's wishes or were able to reassure them. On the other hand, they were motivated to reduce low-value care by a focus on improving patient care.

### Patient factors

Patients' knowledge of the potential harm, lack of benefit and cost of low-value care, facilitated its reduction. For example, when patients with chronic obstructive pulmonary disease were informed in a focus group about the lack of benefit of inhaled corticosteroids, they felt a need to immediately reduce them. However, de-implementation was hindered by frightening stories or incorrect information on the internet. Patients were sometimes afraid of a disease, such as gastric cancer when they had dyspepsia, and wanted reassurance. A lack of trust in, or suspicion of, their clinician also hindered de-implementation.

### Professional interactions

Regarding the professional interactions, de-implementation was hindered by a lack of support and trust, as well as a lack of coordination and collaboration. For example, it was sometimes unclear which clinician was responsible for reducing the low-value care. The convenience and high accessibility of the low-value care also hindered de-implementation. An example of this is the use of standard laboratory packages in the medical ordering system. The growing consciousness among clinicians that more is not always better, as well as good collaboration and support, facilitated de-implementation.

### Incentives and resources

Regarding incentives and resources, de-implementation was hindered by a lack of time, both to communicate with the patient and to participate in the project. It takes more time not to provide low-value care, for example, because patients need to be taught how to check their own skin for cancer in order to reduce follow-up visits to the dermatologist. A potential reduction of revenue was also a barrier to de-implementation in many projects. Clinicians felt hindered to reduce procedures that are reimbursed, such as surveillance visits and insertion of a catheter. In addition, several hospitals and clinicians did not participate in a project because of a fear of reduced revenue.

### Experiences with strategy components

Below, we describe the experiences reported frequently by the project teams, the target clinicians, and the patients

regarding the different components of their de-implementation projects. Online supplemental file 5 shows all experiences.

### Education

Educating clinicians was seen as a useful component of the de-implementation strategy as it enabled them to receive up-to-date information about the low-value care and its side effects. Project 5 included a second educational meeting which focused on practising on a simulated patient, and project 3 showed and discussed a video on communicating with a patient, which helped clinicians to explain to the patient why the care provided is of low value. However, meetings were sometimes either hard to schedule, or could not be attended by all the clinicians. It helped to use existing structures such as weekly meetings. Clinicians found educational material, such as a pocket card, useful. We noted that a lack of repetition contributed to falling back into old patterns. Some terminology, such as 'unnecessary care', and the focus on costs, caused resistance among clinicians.

### Clinical champions

Two projects appointed clinical champions in the participating hospitals. Their task was to bring the subject regularly to the attention of their colleagues and to further spread the educational materials or feedback reports. The way clinical champions fulfilled their role varied. Some spread the messages more actively than others. Clinical champions who left the department or worked in a laboratory did not have as much influence because they did not work near the target group.

### Feedback

Giving feedback to clinicians offered insight into the prevalence of low-value care and comparing their own performance to those of their peers motivated them to perform better. Some clinicians' first reaction was scepticism towards the validity of the data. After reassurance that the data were valid, these clinicians were able to acknowledge that there was room for improvement. Moreover, they were willing to improve. Some projects found the data collection for the feedback time-consuming or even impossible to achieve in time.

### Patient information

Patient information was a valuable de-implementation strategy component, especially in the projects where the patient was an important factor, such as in the reduction of surveillance visits after basal cell carcinoma. However, some factors regarding the spread and content of the material may have limited its effect in other projects. Distribution of the material to patients was not always optimal. Some clinicians considered the information too difficult for patients to understand. Lastly, some clinicians reported that, contrary to its aim, the video clip and poster on vitamin testing in the waiting room led to more requests for vitamin tests, especially for general practices with low preintervention rates of vitamin tests.

### Organisational changes

Organisational improvements in ordering systems or the structure of electronic patient records helped to break habits, although implementing these changes was difficult and took a long time. According to the clinicians, giving routine attention to the subject helped them to remember the message.

### Financial incentives

One project tried to arrange a shared savings contract with insurers, but this could not be achieved within the time frame of the project.

### Project approach

The project leaders reported that they found it very valuable to perform a problem analysis and so achieve greater insight into the context surrounding the practice of low-value care. They used this information to tailor their de-implementation strategy to meet the needs of clinicians and patients and to tackle the barriers that they experience. The problem analysis also created support for the upcoming strategy among the target group. Several project leaders also thought that having a clinician in their project team was essential for recruiting hospitals or general practitioners (GPs) and for providing the education. Lastly, some project leaders found it challenging to collect the right data to evaluate their strategy, because routine hospital or GP data proved to be time-consuming to acquire, was not up to date, or provided insufficient detail to distinguish low-value from high-value care.

## DISCUSSION

### Effects on clinical practice

Five out of the eight projects found a reduction of low-value care following their de-implementation strategy. Two of these five projects compared their results to a control group and found greater reductions in the intervention group. Three out of the eight projects found no effect of the de-implementation strategy. One of these did show a significant reduction in the control group, while another project showed equal reductions in both the control and intervention groups. Both projects reported that the low-value care they targeted received a lot of attention from clinicians nationally, which could have blurred the effect of the strategy and explain the reduction that they found across the country. A comparable dissemination process of seven Choosing Wisely recommendations that recommended against low-value care practices has resulted in a reduction in two out of the seven low-value care practices.<sup>6</sup> This could suggest that dissemination of recommendations including publicity can be sufficient for reducing a part of low-value care practices. The last project with no effect found a non-significant reduction in low-value care in the intervention period, but this was followed by a significant increase in low-value care use after the intervention period, indicating that any potential effect disappeared directly. Unfortunately, this happens more often to de-implementation projects.<sup>26</sup> This shows the

importance of choosing interventions that have sustained results, such as system-focused interventions.<sup>27</sup>

### Barriers and facilitating factors for de-implementation

A lack of time for the patient, an inability to reassure the patients, a desire to meet the patients' wishes and the financial consequences, were frequent barriers to successful de-implementation experienced by clinicians in our study. Both clinicians and patients were hindered by their fear of disease and their search for reassurance, and facilitated by knowledge of the harm associated with low-value care. Reducing low-value care is easier when it is sufficiently supported by the evidence and by consensus among clinicians. Improved collaboration between professions, improved accessibility of the alternative to low-value care and media attention can help to reduce low-value care.

Several of these barriers and facilitators, such as the clinicians' move away from harmful care and their fear of missing a diagnosis, could be connected to the clinicians' motivation to provide the best care for their patients. Two recent studies confirm the importance of harm reduction as a motivator.<sup>12 28</sup> Another connecting theme seems to be the effort that goes into providing less care and communicating this with patients. Patient expectations and a lack of time to turn these around are frequently reported barriers to reducing low-value care.<sup>29–37</sup> In our eight projects, a fear of malpractice was not identified as a barrier, contrary to several other studies from the USA.<sup>29 33</sup> This might indicate that malpractice claims have a smaller influence in the Netherlands. Other studies confirm this. Only 10% of GPs in the Netherlands provide low-value care because of a fear of claims<sup>35</sup> compared with 50%–73% of the primary care physicians in the USA.<sup>33</sup> Fear of malpractice did not emerge at all in our study, possibly because of clinicians' socially desirable responses.

### Experiences with strategy components

Repeated education on the low-value care and on patient communication, as well as feedback were highly valued components of the de-implementation strategies. However, they were hindered by a lack of time to participate in the projects, and difficulties with the availability of data. Patient information was highly valuable when the low-value care was requested by patients. Choosing the right message and content appeared to be crucial for successful patient information.

Two systematic reviews found that multicomponent interventions have the greatest potential in reducing low-value care.<sup>2 8</sup> Two of our projects which targeted only patients achieved significant reductions in low-value care. This suggests that a single intervention can also be effective, although the success of any intervention is generally hard to predict and is likely to depend on the match between barriers and facilitating factors as well as the chosen strategy.<sup>38</sup> Furthermore, Colla and colleagues concluded that supporting clinical decisions, performance feedback and provider education are promising

## Box 2 Practical recommendations for de-implementation projects

Practical recommendations for de-implementation projects based on our evaluation are:

- ⇒ To reduce only low-value care that has sufficient evidence, and consensus among clinicians, of being of low value. When the field is not ready for de-implementation, you risk provoking discussions among clinicians, achieving less or no effect.
- ⇒ To perform a problem analysis of the low-value care practice you are aiming to reduce and study the context of your project. Then tailor the de-implementation strategy to the barriers and facilitating factors you have found.

Some tips about specific parts of the strategy are:

- ⇒ Educating clinicians and improving their communication skills can be useful, especially when existing meetings are used and the message is repeated.
- ⇒ To provide regular feedback if data are easily available in order to motivate clinicians to reduce their use of low-value care.
- ⇒ To provide information material for patients when they request the low-value care, while ensuring it is the right length, has the right message and is distributed by clinicians.
- ⇒ To promote organisational changes such as providing tools to support clinical decision-making in order to challenge previous patterns of practice.
- ⇒ To be aware that a lack of time and a loss of revenue can be major barriers to de-implementation. There may be no easy solution for this.
- ⇒ To focus on improving the quality and safety of care instead of saving costs. Clinicians and patients are motivated to reduce low-value care when they learn about its burden and harm.
- ⇒ To be aware that reducing low-value care can evoke fear and uncertainty in both clinicians and patients.

strategies.<sup>2</sup> Our study confirms this while adding patient information as another promising strategy. Additionally, in our practical recommendations (box 2), we provide conditions for the success of these strategies.

Our study is the first that combines the lessons from multiple multicentre de-implementation projects. It is complementary to the study by Stinnett-Donnelly and colleagues that described the lessons from local de-implementation projects in one medical centre.<sup>11</sup> They found that the value of a project, such as the reduction in patient harm, promotes de-implementation. They also showed that more controversial care practices among clinicians require more effort to de-implement, and that data collection could be labour-intensive.<sup>11</sup> Parker and colleagues identified several challenges and facilitators for leaders of de-implementation projects, such as the availability of data and harm reduction.<sup>12</sup> We confirmed their findings and identified more lessons regarding both the barriers and facilitating factors, and the promising components of a de-implementation project.

### Strengths and limitations

The strength of our study is the prospective design, which enabled us to observe the project leaders' experiences throughout all steps of the projects. Another strength is

that we were able to combine their experiences since the projects had the same structure, even though they were performed in different regions and targeted different practices. However, this diversity can also be a limitation with regard to their comparability.

The validity of our results depends on the quality of the methodology used in the eight projects. Three of the five projects that achieved a reduction in low-value care did not compare their intervention to a concurrent control group. Before that reason we do not know to what extent their reduction in low-value care can be attributed to a national trend instead of to the de-implementation strategy adopted by the project. It could therefore be the case that our results overestimate the effects of a de-implementation strategy. Other items that indicate the quality of a project, such as blinding or randomisation, are reported in the papers of the individual projects (referenced in online supplemental file 3).

A second limitation is that the qualitative analysis of the projects is conducted by two authors who were part of the coordinating team that supported the eight projects. This might have biased both the experiences that the project leaders reported in the interviews and logs, and the authors in their analysis, to present a more favourable picture of the projects.

The projects' method and time point of identifying the barriers and factors facilitating de-implementation varied. It is possible that some projects missed relevant factors. Regarding the experiences with the different components of the projects, the results are based on the evaluation and subjective experiences of the project leaders. Other project leaders may have different experiences.

### Implications for research and practice

Many hospitals and general practices in the Netherlands participated in the eight projects described. This has amounted to the prevention of tens of unnecessary endoscopies and dermatology visits, hundreds of unnecessary catheters, and thousands of unnecessary vitamin and laboratory tests. The next step is to sustain these results and spread them to other hospitals in the Netherlands. The five successful projects are currently being spread throughout the Netherlands and of three projects the long-term effects will be measured. The changes that our projects achieved should transcend their project setting and become a permanent part of clinical practice. However, few de-implementation projects evaluate long-term sustainability and more knowledge on this is required.<sup>3</sup> The majority of the literature on the spread and dissemination of projects is focused on implementation rather than de-implementation, such as the theory of Rogers.<sup>39</sup> Research is necessary to evaluate whether these theories are also relevant for de-implementation projects.

The costs saved to Dutch society associated with a reduction in low-value care are hard to achieve and measure. Some savings can only be realised by reducing equipment and personnel, which is hard to realise in the short term. Also, the costs associated with all potential unintended



consequences of the strategy, such as an increase in the use of other care, should be monitored. Further research is necessary into the potential for cost savings.

Our findings can support clinicians and researchers in leading more successful de-implementation initiatives by providing examples of the barriers, facilitating factors and valuable components drawn from our eight de-implementation projects. We have combined their results and experiences and translated them into practical recommendations for de-implementation projects (box 2).

## CONCLUSIONS

Successfully reducing low-value care is possible in spite of the powerful barriers opposing it. The eight de-implementation projects managed to recruit many hospitals and general practices. Five of these achieved significant results without measuring negative consequences. We offer practical recommendations for reducing low-value care successfully and preventing patient harm. These include: reduce only low-value care that is supported by sufficient evidence; tailor the strategy to counter the barriers; use repeated education and feedback for clinicians; provide carefully developed patient information when patients request the low-value care; and adapt the organisation to support this change.

**Acknowledgements** The authors thank the independent committee that selected the eight projects, the advisory board that advised on the programme, and editor Tony Sheldon who helped to create this manuscript, for their time and effort.

**Collaborators** To do or not to do programme collaborators: Corina de Jong, Janwillem Kocks, Aniek de Coninck, Harry C Schouten, Leti van Bodegom-Vos, Tessa Rietbergen, Bart J Laan, Suzanne E Geerlings, Evelien IT de Schepper, Saskia F van Vugt, Prabath WB Nanayakkara, Renuka S Bindraban, Sven van Egmond, Marlies Wakkee, Judith J de Jong, Joost PH Drenth.

**Contributors** EWW, SAVD, GPW, LH, PH and RBK were involved in the study concept and design. RBK obtained funding. EWW and PH performed the interviews. EWW performed the analyses, which PH verified. EWW, SAVD, PH and RBK interpreted the data. EWW drafted the manuscript. RBK was the guarantor of this study. The To do or not to do programme collaborators led the eight projects that were evaluated. All authors participated in critical revision of the manuscript for important intellectual content. The corresponding author attests that all listed authors meet authorship criteria and that no others meeting the criteria have been omitted. All authors approved the final version of this paper and agree to be accountable for all aspects of the work. The To do or not to do programme collaborators group consists of the following authors: Corina de Jong, Janwillem Kocks, Aniek de Coninck, Harry C Schouten, Leti van Bodegom-Vos, Tessa Rietbergen, Bart J Laan, Suzanne E Geerlings, Evelien IT de Schepper, Saskia F van Vugt, Prabath WB Nanayakkara, Renuka S Bindraban, Sven van Egmond, Marlies Wakkee, Judith J de Jong, Joost PH Drenth. The To do or not to do programme collaborators led the eight projects that were evaluated. They recruited the participants, performed the problem analysis, the strategy, and the evaluation of their project, and they were interviewed for this study. Corina de Jong and Janwillem Kocks led project 1; Aniek de Coninck and Harry C Schouten led project 2; Leti van Bodegom-Vos and Tessa Rietbergen led project 3; Bart J Laan and Suzanne E Geerlings led project 4; Evelien IT de Schepper and Saskia F van Vugt led project 5; Prabath WB Nanayakkara and Renuka S Bindraban led project 6; Sven van Egmond and Marlies Wakkee led project 7; and Judith J de Jong and Joost PH Drenth led project 8.

**Funding** This work was supported by ZonMw, the Netherlands Organisation for Health Research and Development (grant number 839201 002).

**Competing interests** All authors had financial support from ZonMw for the submitted work. ZonMw is an independent organisation for health research and development that distributes grants on behalf of the Dutch government. Kocks reports grants and personal fees from AstraZeneca, grants and personal fees from

Boehringer Ingelheim, grants from Chiesi, grants and personal fees from GSK, grants from Mundi Pharma, grants from TEVA, outside the submitted work; all fees go to the institute. Drenth reports grants from Gilead outside the submitted work; all the proceeds go directly to the institute.

**Patient and public involvement** Patients and/or the public were involved in the design, or conduct, or reporting, or dissemination plans of this research. Refer to the Methods section for further details.

**Patient consent for publication** Not applicable.

**Ethics approval** This study involves human participants. For this interview study that evaluated the effects and experiences of eight projects, ethical approval was not required under Dutch national law. The eight projects that are studied obtained ethical approval before the start of their study. Participants gave informed consent to participate in the study before taking part.

**Provenance and peer review** Not commissioned; externally peer reviewed.

**Data availability statement** Data are available upon reasonable request. De-identified reports of the eight projects are available (in Dutch) from the corresponding author upon reasonable request.

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