Using a national archive of patient experience narratives to promote local patient-centred quality improvement: an ethnographic process evaluation of ‘accelerated’ Experience-based Co-design

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Using a national archive of patient experience narratives to promote local patient-centred quality improvement: an ethnographic process evaluation of ‘accelerated’ Experience-based Co-design

Abstract

Objectives: To evaluate an accelerated form of Experience-based Co-design (EBCD), a participatory action research in which patients and staff work together to improve quality; to observe how acceleration affected the process and outcomes of the intervention.

Methods: An ethnographic process evaluation of an adapted form of EBCD was conducted, including observations, interviews, questionnaires and documentary analysis. Whilst retaining all components of EBCD, the adapted approach replaced local patient interviews with secondary analysis of a national archive of patient experience narratives to create national trigger films; shortened the timeframe; and employed local improvement facilitators. It was tested in intensive care and lung cancer in two English National Health Service (NHS) hospitals. A total of 96 clinical staff (primarily nursing and medical), and 63 patients and family members participated in co-design activities.

Results: The accelerated approach proved acceptable to staff and patients; using films
of national rather than local narratives did not adversely affect local NHS staff engagement, and may have made the process less threatening or challenging. Local patients felt the national films generally reflected important themes although a minority felt they were more negative than their own experience. However, they served their purpose of ‘triggering’ discussion between patients and staff, and the resulting 48 co-design (improvement) activities across the four pathways were similar to those in EBCD but achieved more quickly and at lower cost.

Conclusions: Accelerated EBCD offers a rigorous and relatively cost-effective patient-centred quality improvement approach.

Preferred keywords: experience-based co-design, narrative, quality improvement, patient experience
Background

Measuring or improving patient experience?

Measuring, understanding and improving patient experience is a priority for healthcare systems worldwide\(^{(1)}\). Policy documents in several countries describe ‘patient experience’ as a core component of care quality, drawing on frameworks for patient-centred care developed by, for example, the Picker Institute (following Gerteis et al.\(^{(2)}\)) and the US Institute of Medicine\(^{(3)}\). The English NHS has led the way in measuring patient experience by introducing the first nationally mandated patient survey\(^{(4)}\).

Justifications for seeking to improve patient experience include moral humanitarian concern; the need to maintain political legitimacy in state-funded systems; and the need to increase/maintain market share in more competitive systems. A growing international body of evidence suggests patient experience, safety and clinical effectiveness are linked, showing positive associations between patient experience measures and both self-rated and objectively measured health outcomes\(^{(5, 6)}\) as well as positive organisational outcomes\(^{(7-13)}\).

Despite the strong policy focus, some patients continue to report unsatisfactory experience in even the best funded health systems\(^{(14)}\). In England, recent
government policy has identified the lack of a genuinely patient-centred approach.

Examples of poor care have dominated healthcare debate, the Francis Inquiry into Mid Staffordshire Hospital being the most widely reported\(^{15}\).

Robert and Cornwell\(^{4}\) argue that the focus on experience surveys and targets may paradoxically have ‘contributed to a tick-box or compliance mentality on the part of [hospital] management boards… [and] lulled them into thinking that they were paying attention to their patients’ experiences’. Experience data presented to NHS hospital boards is often ‘noted for information’ rather than leading to action points and organisational learning\(^{16}\).

Failure to attend to basic human needs and dignity, as documented in the Francis Report, has resulted in numerous recommendations to re-educate healthcare staff and initiatives to improve compassion. No-one would dispute the importance of compassion and kindness, but the danger is that the energy behind this latest series of initiatives may produce more targets, tools, metrics and inspection, reinforcing the ‘compliance mentality’ as organisations work to measure how compassionate their staff are, rather than the more fundamental task of focussing on patients and
changing the culture to enable this to happen.

A large, growing body of quantitative and qualitative research\(^{[3, 17-19]}\) already identifies aspects of care that patients and families think are important. The challenge is to find meaningful ways to enable organisations to learn from this evidence and move beyond data-gathering to using patient experiences to improve care\(^{[20-22]}\). Surveys provide useful information to make comparisons between sites and across time, but as a recent review of the impact of the UK National Inpatient Survey\(^{[23]}\) concludes, ‘the inpatient survey is not in itself a quality improvement tool’ and ‘simply providing hospitals with patient feedback does not automatically have a positive effect on quality standards’.

There is increasing interest in the use of in-depth qualitative research to gain richer, meaningful accounts of what being a patient is like, yet healthcare providers struggle to use qualitative experiential evidence for improvement\(^{[16]}\).

**The Value of Narrative and Experience-Based Co-Design (EBCD)**

Narrative persuasion is an established psychological theory\(^{[24]}\) demonstrating that narrative is a powerful way of accessing human experience, transporting us to another world and enabling us to see things through another’s eyes. Narratives can engage care providers, at a deep emotional level, in reflecting on how services could be
improved \cite{25, 26}. Narratives are not gathered because they are assumed to be objective, accurate or verifiable but because they are uniquely human and subjective, describing not a fact or a reality but a recalled experience or set of experiences.

Experience-Based Co-Design (EBCD), a participatory action research approach, marks a significant contribution to using narratives and involving patients in quality improvement in healthcare \cite{26}. The approach has been implemented in over 60 services internationally and independently evaluated in both England and Australia \cite{27-29}. It is effective in making specific improvements to particular services, as well as wider improvements within - and sometimes between – healthcare organisations.

EBCD projects typically last 12 months, beginning with a 6-month ‘discovery’ phase, in which local patients and staff are interviewed about their experiences of a service. Patient narratives are video-recorded, and used to develop a ‘trigger film’ to stimulate discussion between staff and patients about potential quality improvements. The discovery phase is characterised by rigorous, narrative-based research with a broad sample, rather than relying on a few anecdotes. Equally important is the subsequent co-design phase, in which patients, families and staff join as partners in small working groups to set priorities for quality improvement, and design and implement changes.
However, the discovery phase is felt by staff to be lengthy and costly, and has been reported as a barrier to adoption of the approach. Undertaking 5-6 months of qualitative interviewing on each pathway in each hospital is seen as impractical.

‘Accelerated’ EBCD and our research questions

The Health Experiences Research Group (HERG) at the University of Oxford collects and analyses video/audio-recorded interviews with people about experiences of illness. It has an archive of around 3000 interviews, covering over 80 different conditions or topics. Selected interview extracts are disseminated for a lay audience on www.healthtalkonline.org. In this study we investigated whether this archive could replace the need for local discovery interviews, thus making EBCD faster and cheaper; we describe this as accelerated EBCD or ‘AEBCD’. By using national rather than local patient interviews, we aimed to halve the cycle from 12 to 6 months. Our research questions can be summarised as:

1) What improvement activities does the approach stimulate (for what cost) and how do these compare with traditional EBCD?

2) How does using films of national rather than local narratives affect the level and quality of engagement with service improvement by local NHS staff?

3) How well do local patients feel national narratives represent themes important to
their own experience?

Methods

The intervention was an adapted form of EBCD: we used national trigger films, shortened the time frame and employed local improvement facilitators (senior nurses or service managers in the participating hospitals who were trained in EBCD). It was tested in intensive care and lung cancer in two English NHS hospitals. 96 staff (primarily nursing and medical), and 63 patients and family members participated.

The trigger films were derived from secondary analysis of the Oxford University collections of patient experience interviews. We re-analysed all the interviews in the archive with people with lung cancer, and individuals and family members with experiences of intensive care. The local facilitators interviewed staff to understand their experience of providing care. Thereafter the process followed the usual EBCD pattern: the film was shown to local patients in a workshop; staff had a separate meeting to discuss the results of their own feedback. Staff and patients then came together in a further workshop to view the film, agree priorities for quality improvement and establish co-design working groups to take these priorities forward.

An ethnographic process evaluation using multiple data sources was conducted to observe implementation in both pathways in each hospital. Ethnography can
contribute to process evaluation by providing rich accounts of activities, projects and programmes (31). The ethnographer was in post throughout and was therefore able to observe all stages of the project. Data collection took place between November 2011 and December 2012 including 155 hours of observations (of facilitator training sessions, staff and patient workshops, joint events, co-design meetings and celebration events, project steering group and core group meetings); 30 interviews (designed around the following themes: involvement in the project and perceptions of the process; project contribution to service delivery; and project sustainability and legacy); two group interviews with four patient participants to discuss their involvement and perceptions of the process during each step of the intervention; cost analysis; and four service improvement logs (tables listing local improvement activities completed by the facilitators). Interviews, training sessions and events were audio-recorded and transcribed. Observations and brief conversations were recorded as fieldnotes and 22 reflective diaries were completed (standard forms e-mailed to project team members on a monthly basis to gather direct information and reflections on issues and concerns regarding the day-to-day running of the project). 166 evaluation questionnaires (self-completion questionnaires, identical to those used in previous EBCD projects) were distributed to all participants at the end of each event and collected by the evaluator.

Reports from 12 previous EBCD projects were analysed for comparison.
Transcripts, documents, fieldnotes and emails were entered into NVivo. Coding was based on our research questions (see above), as well as emerging themes (such as different styles of facilitation and implementation). Data were tabulated using framework analysis, a method of drawing up summarised data by theme and by case into a matrix (32). Data analysis involved the following stages: familiarisation with AEBCD and EBCD data; thematic analysis of qualitative data; tabulation and graphical representation of quantitative questionnaire and quality improvement data; indexing and developing a comparative framework based on key themes. Members of the project team refined the analysis at a two-day workshop.

Results

The accelerated approach proved readily acceptable to staff and patients; using films of national rather than local narratives did not adversely affect local staff engagement, and may have made the process less threatening or challenging. Local patients felt the national films generally reflected important themes although a minority felt they were more negative than their own experience. However, they served their purpose of ‘triggering’ discussion and the resulting improvement activities were similar to those in EBCD, but this was achieved more quickly and at reduced cost.

Improvement activities and cost

We recorded 48 co-design activities across the four pathways, and these were similar
in nature and scale to those typically seen in EBCD (see Box 1).

Although small-scale changes and process redesign within one service area are the most common result of both AEBCD and EBCD, it was observed that small-scale change is often remarkably complex to implement, and what looks like a small change can be immensely valuable to patients. Moore and Buchanan have recently referred to this as ‘sweating the small stuff’ (33). For example ensuring an inpatient’s belongings follow them as they move around the hospital seems a minor issue, but as the facilitator involved in this change noted, being without your false teeth when talking to the consultant is both practically difficult and humiliating.

Cost reductions were (as anticipated) in the ‘discovery’ rather than the ‘co-design’ phase; in AEBCD the cost of developing a trigger film was £8,289, compared to £30,485 for EBCD (see Box 2). In AEBCD this is a one-off cost so future projects would only incur the co-design phase costs, which in this project were £20,276 (over half of which was for facilitator salary). Full details of costings are reported elsewhere (34).

Acceptability and impact - staff participants
Previous evaluations have EBCD to be highly acceptable\textsuperscript{(27, 28, 35)}. Before the study, we speculated that removing local patient interviews might negatively affect local ownership and engagement. In fact, the project was positively received by both patients and staff; results from post-workshop questionnaires suggested broadly similar satisfaction levels to traditional EBCD. The interviews and comments recorded during observations featured many positive accounts. The opportunity to hear directly from patients and carers had a transforming effect on some staff. One described herself as ‘a better nurse because of it.’ A facilitator commented:

\textit{The absolute core of it is the humanistic connections. Because of hearing patients’, relatives’ and staff experience it acts as a catalyst and gives you energy to keep going and make the change, and make sure it happens [...]}

\textit{So many people have echoed that without me saying anything. I have not had to convert people into this methodology now. People just get it and wonder why we’ve never done it before. (Facilitator, interview)}

It was striking that the approach successfully engaged medical staff, a group often not closely involved in quality improvement projects\textsuperscript{(36)}. One consultant reflected how the project gave staff an opportunity to see the person behind the patient:
I think [AEBCD] is phenomenal [….] I don’t have any idea, as a unit we don’t
have an idea of what is beyond what I’m seeing. This informs that process. So I
can see that this person is not only a human being, but he is also a father, he is
a son, he is a brother, he is a friend, he is a cousin, he’s a plumber or an
electrician, he is a sportsman, he has an interest in horse-riding, whatever it
happens to be. He has a dog, he has a budgie, he has plans, he has
expectations, he has regrets, he has feelings.’ (Consultant, intensive care,
interview)

A nurse reported her changed attitude immediately after the first co-design workshop:

‘I have already changed the way I think and care for patients even though we
haven’t started implementing changes yet. I have a better understanding now
of how things are from the patients’ perspective.’ (Nurse, intensive care,
fieldnote of conversation)

Our data suggest face-to-face encounters with patients in co-design groups often had a
profound effect on staff in making them think differently about their practice and
reconnect with their core professional values, resulting in renewed motivation. A
senior manager reported that a staff member in intensive care had said it was the first
time in 20 years of practice that he had sat down and talked to patients in this way. A
senior lung cancer nurse (unprompted) told a meeting of the project advisory group
that it was the most inspiring thing in her professional career.

Critical aspects of the process included creating space to think and listen to others
(both patients and colleagues):

‘I think the most important things were that staff really appreciated the time to
think about the experience. It became apparent that they perhaps didn’t have
or make time to reflect on what they do in their daily workings, so I think they
actually found it quite cathartic and therapeutic. […] I think the staff really
appreciated that their point of view was being listened to because up until that
point the political drive had always been patient experience and now all of a
sudden we were interested in staff experience.’ (Facilitator, interview)

One principle of EBCD is that seeing one’s own patients talking about their experiences
will be an especially powerful insight that motivates staff to rethink how care is
provided. We were concerned that using films of national rather than local narratives
might enable staff to distance themselves from the experiences shown and claim such
things do not happen in their trust. Staff were therefore asked to rank their experience
of seeing the national trigger film at the joint patient-staff co-design workshops. They
consistently ranked the film and its representation of experiences more highly than 
patients at the same events, always ranking it as good or excellent. It was observed in 
one site that staff defended the film when patients said they felt some aspects were 
too negative and therefore not an accurate reflection of care in the trust (see below); 
staff felt able to challenge this perception, confirming that such things did indeed 
happen locally on occasion and sharing experiences of witnessing sub-optimal care.

The observational data confirmed that the film achieved the goal of triggering thought, 
discussion and plans for action, and that using national narratives seemed to allow 
staff to engage comfortably with patient experiences.

One facilitator reflected that the use of national narratives may have helped staff feel 
less threatened by negative comments and able to externalise criticisms of care.

It made me wonder that [...] if the DVD had been produced locally, and we were 
all in a room together, and that was maybe the first time that healthcare 
professionals had had feedback from patients, as to how potentially devastating 
that could have been [...] It did make me grateful that I didn’t have to deal with a 
potential fallout if a patient had criticised a member of staff. [...] I think that 
potentially could have quite catastrophic effects. That’s why I wondered whether
the Healthtalkonline actually minimised that risk.

It is arguable that a degree of unsettlement amongst staff can be productive (36), and ensure opportunities for deep emotional reflection and self-challenge. At the same time, a genuine staff-patient coalition is central to EBCD, and may be easier to achieve if staff do not feel personally criticised.

Acceptability and impact – patient/carer participants

Intriguingly, patients and relatives sometimes felt the national films were more negative than their own experiences; this may be partly because people who volunteer for such projects feel positive about their local hospital and want to give something back for potentially life-saving care. However, the great majority of those present rated the film as ‘good’ (71%) or ‘excellent’ (14%) in answer to the questionnaire item 'Seeing the film today - did you think it was a good representation of what it is like to be a [lung cancer/intensive care] patient at X NHS Trust?’ A few (8.5%) rated it as ‘average’ but no-one as a ‘poor’ representation of their experience. Data were incomplete for 6.5%.

Workshop observations showed that after discussing the film, patients in both
pathways gradually revealed more things they felt had not gone well in their care, and
began to reflect on aspects of care which could be improved. It thus seems to have
served a similar role to the film in other EBCD projects – triggering discussion. Overall
reflections on the process focus on the value of meeting with other patients/relatives
and staff and getting truly involved in change.

‘It was great to see you all at the “Patient Experience” event this week, what a
wonderful thing it will be if we can contribute to and change what is already an
incredible service that the intensive care department already provide - it makes
me feel humble to be part of such a cause. Sorry to be gushing, but [facilitator]
you truly are a guardian angel. I also feel that this experience should be shared
with other hospitals as it seems they could learn a lot.’ (Patient, intensive care,
email correspondence)

A lung cancer patient contrasted AEBCD with normal consultation in which ‘people say
I’ll do this, I’ll do that, and nothing happens’. An intensive care participant said, ‘I feel I
was really listened to’ and it was ‘much better than expected.’ Several expressed
surprise that they had felt able to contribute as equal partners and that their views
were taken seriously.

‘I wondered what, if anything, will be taken on board. To be honest with you,
everything has been taken on board, and that in itself was a complete surprise.’

(Patient interview)

Patients who presented at celebration events expressed strong support for the process. The ethnographer noted:

‘There was a comment from a patient within the event who said that everything that had been discussed had been taken forward into improvements... ‘our views were not dismissed, they were looked at and things changed’. ‘Thank you for looking at the little things.’ (Excerpt from ethnographer’s field notes at celebration event)

Conclusions - the fundamental importance of co-design

This study was designed to answer specific questions about whether local patient discovery interviews in traditional EBCD could be replaced with a nationally derived trigger film, how this would affect patient and staff engagement, and whether this would affect the resulting quality improvement activities. For this study, we reanalysed just three of the interview collections in the archive. At the time of writing, it contains collections on over 80 different conditions, with more added each year. Sister projects have also been established in eight other countries under the auspices of the DIPEx International collaboration (see www.dipexinternational.org). This is an
enormous potential resource for new films on a range of topics.

As the intervention progressed, it became increasingly apparent that agreement or identification with the film content was not central. The purpose of the films in both AEBCD and traditional EBCD is simply to ‘trigger’ discussion and enable people to share ideas and concerns; then joint work to redesign care can begin. If national trigger films are ‘good enough’ to initiate such conversations, local specifics can be brought into discussion along the way, and even disagreement with the content can generate productive discussion. But the element of the intervention that left both staff and patients feeling energised and empowered is the direct mutual encounter, the active partnership in co-design groups to achieve change, and the sense of tangible results.

A realist review of studies using participatory research argues that the link between process and outcome in participatory research can best be explained by the middle range theory of ‘partnership synergy’; that is to say, ‘multiple stakeholder collaboration creates or enhances research outcomes beyond what could be achieved by a single person or organization working under similar conditions’. While on occasion the result can be conflict and negative outcomes, the review demonstrates that partnership synergy brings many benefits, including culturally and logistically appropriate research; better quality of outputs and outcomes over time; increased
1. sustainability of project goals beyond funded timeframes; system changes; and new,
2. unanticipated projects and activities. This resonates strongly with our findings, and is
3. the antithesis of the problems underlying the Francis Report of failing to listen to
4. patients or take their concerns seriously.

5. It is possible to implement improvements based on patients’ experiences without
6. involving them in the change process. Staff could be shown a trigger film or interview
7. data previously collected from patients and then decide as a staff working group what
8. needs doing. It is true that just seeing patient narratives on film can in itself have a
9. powerful effect. But our observations suggest face-to-face encounters with patients
10. have been even more transformative, inspiring and revelatory to staff in making them
11. think differently about their values and practice. Having continued patient involvement
12. helps ensure improvements really do address patient concerns and holds staff to
13. account to see change through. Patients’ physical presence constantly reminds
14. everyone who change is for, and why it matters compared to other potentially
15. overwhelming work pressures and demands. When staff report that this is the first
16. time in 20 years that they have really talked to patients in this way or that it is the
17. most rewarding thing in their careers, the potential of EBCD to reconnect staff with
18. their fundamental values of care and compassion is striking. Patients, too, report a
19. new level of appreciation for staff, a belief that they will be listened to and that change
is possible, and renewed trust in their local NHS.

There is limited evidence on the costs and cost-effectiveness of patient-centred quality improvement. In comparing EBCD and AEBCD, this study has for the first time analysed the costs of EBCD. The next stage is to understand more about relative cost-effectiveness as no quality improvement activity is cost-free; it requires staff time, energy and commitment to make it happen. We believe there may still be scope for further economies in developing trigger films, whilst ensuring they remain based on high quality research into patient experience.

As ever with in-depth qualitative research, our findings relate to a small number of sites and may not transfer easily to other contexts; further evaluation would be helpful. What we have demonstrated, however, is one way to implement a proven technique faster and more cheaply without undermining its effectiveness. But, perhaps more importantly, we believe it is an approach that can revitalise staff. This is supported by work demonstrating that staff experience and patient experience are closely linked (39), providing a new way of stimulating compassionate care.


23. DeCourcy A, West E, Barron D. The National Adult Inpatient Survey conducted in the English National Health Service from 2002 to 2009: how have the data been used and what do we know as a result? BMC health services research. 2012;12:71.


Box 1 Comparison of improvement activities in AEBCD and EBCD

Changes in the four AEBCD pathways were compared with those in two previous EBCD projects on breast cancer and lung cancer. The scale of the changes was similar. There were 28 activities across the two EBCD pathways, compared to 48 across the four AEBCD examples. There were similarities in terms of the co-design group topics. For example, two EBCD lung cancer groups focused on the same overall topics as the AEBCD lung cancer groups: information and diagnosis giving. There are also identical improvement actions in the EBCD and AEBCD lists. For example, name boards were introduced so that patients would know staff names and both groups identified the importance of a special, private room for diagnosis giving. Finally, there was a similar distribution of activities, with more small scale changes and process redesign within teams than wider process redesign between services and between organisations. An overview of the number and nature of changes in EBCD and AEBCD is given below:

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<th>Small scale changes</th>
<th>Process redesign with teams</th>
<th>Process redesign between service activities</th>
<th>Process redesign between organisations</th>
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<td><strong>EBCD</strong></td>
<td>12</td>
<td>12</td>
<td>2</td>
<td>2</td>
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<tr>
<td><strong>Examples</strong></td>
<td>reviewing and improving patient information; regular updates on waiting times for patients in clinic</td>
<td>designated phlebotomist to reduce waiting time for blood tests</td>
<td>physiotherapists reviewed timing to give patients advice about exercise</td>
<td>link nurse scheme to improve cross-site working and cross-site visibility of test results</td>
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<tr>
<td><strong>AEBCD</strong></td>
<td>21</td>
<td>21</td>
<td>5</td>
<td>1</td>
</tr>
<tr>
<td><strong>Examples</strong></td>
<td>sourcing clocks to aid patient orientation in ICU; more comfortable V-shaped pillows for post-operative patients</td>
<td>new private room identified for receiving support after diagnosis; introducing mini ‘Schwartz rounds’ on ICU</td>
<td>changed process for porters to remove waste avoiding ICU rest times; redesigned discharge summary with input from all professions</td>
<td>improved cross-site information booklet for patients transferring to another hospital for surgery</td>
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Box 2 – Comparison of costs between AEBCD and EBCD

<table>
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<th>Activity</th>
<th>EBCD per pathway (based on the budget for a previous study funded by the Guy’s and St Thomas’ Charity)</th>
<th>AEBCD per new pathway</th>
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<td><strong>Fieldwork to collect patient narratives &amp; staff interviews (and non-participant observation)</strong></td>
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<td></td>
<td>- Researcher travel &amp; subsistence</td>
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<td></td>
<td>- Transcription costs</td>
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<td></td>
<td></td>
<td>Not applicable</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0</td>
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<tr>
<td><strong>Production of ‘trigger’ films</strong></td>
<td>- Editing film</td>
<td>1100</td>
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<tr>
<td></td>
<td>- Researcher time</td>
<td>5289</td>
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<tr>
<td></td>
<td>- Editing</td>
<td>3000</td>
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<tr>
<td><strong>TOTAL</strong></td>
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