Immediate Benefit of Art on Pain and Well-Being in Community-Dwelling Patients with Mild Alzheimer’s

Elodie Pongan, MSc1,2,3, Floriane Delphin-Combe, MSc2,3, Pierre Krolak-Salmon, MD, PhD2,3,4, Yohana Leveque, PhD4,5, Barbara Tillmann, PhD4,5, Romain Bachelet, MSc2,3, Jean-Claude Getenet, MD1, Nicolas Auguste, MD6, Béatrice Trombert, MD, PhD7,8, Jean-Michel Dorey, MD9, Bernard Laurent, MD, PhD1, and Isabelle Rouch, MD, PhD1,2,3

Abstract

Objective: The present report aims to evaluate whether singing intervention can bring an immediate benefit that is greater than the one provided by painting intervention on pain and well-being. Methods: Fifty-nine mild patients with Alzheimer disease were randomized to a 12-week singing (n = 31) or painting group (n = 28). In the present analysis, the immediate evolution of pain and well-being was compared across sessions between the 2 groups using mixed-effects models. Results: We observed a significant improvement in well-being for both singing and painting groups immediately after sessions, compared to the assessment before the sessions. We did not observe this improvement across the sessions for pain intensity measurement. Discussion: Our results revealed that both painting and singing interventions provide an immediate benefit on the patients’ well-being.

Keywords
Alzheimer disease, music, pain, well-being, quality of life

Introduction

Alzheimer disease (AD) is a common disorder that principally concerns patients of 65 years and older,1 with two-thirds of them being aged 75 years and older.2 Patients with Alzheimer disease and related disorders (ADRDs) are more at risk of having physical pain because of the links between pain and cognition3 but also because of their age and their frailty. In fact, these patients with AD often present multiple comorbidities that can induce chronic pain.4 Pain in elderly is more persistent compared to younger patients. In a population of patients with ADRD at a moderate stage, a functional magnetic resonance imaging study showed a higher response to pain during a mechanical stimulation compared to control participants.5 Otherwise, there is ample evidence that pain affects well-being. Pain interferes with activities of daily living, causing emotional distress and impacting psychological health.6 So, this can accelerate the loss of autonomy associated with neurodegenerative disease per se. However, analgesic medications

1 Neurology Unit, CM2R, University Hospital of Saint-Etienne, Saint-Etienne, France
2 Clinical and Research Memory Center of Lyon, University Hospital of Lyon, Lyon, France
3 Institute of Aging I-Vie, University Hospital of Lyon, Lyon, France
4 Institute of Aging I-Vie, University Hospital of Lyon, Lyon, France
5 Lyon Neuroscience Research Center, CNRS, UMR5292 INSERM, U1028, Psychoacoustic and Auditory Cognition Team, Lyon, France
6 Geriatrics Unit, CM2R, University Hospital of Saint-Etienne, Saint-Etienne, France
7 Public Health and Medical Information Unit, University Hospital of Saint-Etienne, Saint-Etienne, France
8 SNA (EA SNA EPIS 4607, Autonomic Nervous System), University of Lyon, Saint-Etienne, France
9 Psychiatry Unit, Centre hospitalier le Vinatier, Lyon, France

Corresponding Author:
Elodie Pongan, MSc, Centre Mémoire de Ressources et de Recherche, Neurology Unit, University Hospital of Saint-Etienne, Saint-Etienne 42055, France.
Email: elodie.pongan@chu-lyon.fr
are not free of side effects, and they can increase the risk of confusion, falls, and cognitive decline. In addition, they can induce a physical and psychological dependence as well as difficulties to withdrawal.2

The French Health Authority recommends combining non-drug therapies with standard treatments. Among them, musical intervention (MI) has already proven usefulness and effectiveness in certain domains such as affective symptoms8 and cognition.9,11 In elderly patients, beneficial effects of MI have also been reported for chronic pain.12,13 However, to our knowledge, the potential effects of MI on chronic pain have never been studied in elderly patients with AD.

The LACMe study aimed to determine the efficiency of singing intervention (SI) versus another artistic activity, namely painting intervention (PI) on chronic pain, mood, and cognition in patients with AD with moderate to severe chronic pain. Quality of life (QoL) was also measured as it is a core marker for the benefit of therapeutic strategies.14,15

In a previous work,16 we have reported the long-term results of the LACMe study, notably showing that after 12 weeks of either SI or PI, patients with mild AD had reduced chronic pain, improved mood, and QoL. Differential effects were shown with better memory performance with the SI and an improvement of depression with the PI, but there was no difference between the SI and PI for pain and QoL. However, these results did not provide information about the immediate benefit of these activities, leading us to report here the more detailed findings after each session.

The potential anaggesic effect of music has been previously reported: The pleasurable character of the music-induced emotions can reduce the painful sensation due to an experimental stimulus.17,18 Pleasant versus unpleasant music can modulate the activity of limbic areas and areas related to the reward system,19,20 which are involved in painful perception as well as well-being. In addition, singing involves breathing and has an effect on endorphin production.21 Finally, music can produce effects on heart rate and breathing rate depending on the tempo and then induces relaxation, mostly during pauses or slower rhythms.22 Singing intervention could therefore have an immediate relaxing transitory power that cannot be observed after a long time period from activities.

The present report was aimed to evaluate with an explanatory design whether SI can bring an immediate benefit that is greater than the one provided by PI on pain and well-being.

Materials and Methods

Study Design

The present analysis was based on data from the LACMe study, a multicenter randomized clinical trial with 2 parallel arms and a blinded end point evaluation. The LACMe design and global results about pre- and postintervention changes were disclosed in 2 previous papers.16,23 This trial was registered at clinicaltrials.gov. (Identifier: NCT02670993).

Population

Participants of the LACMe study were recruited from 3 memory clinics in France. The inclusion criteria were as follows: 60 years and older and probable AD at mild stage, according to the Diagnostic and Statistical Manual of Mental Disorders (Fifth Edition)24 and the National Institute of Neurological and Communicative Disorders and Stroke/Alzheimer’s Disease and Related Disorders Association criteria.25 The severity of cognitive disorders was assessed with the Mini-Mental State Examination (MMSE),26 and only participants with an MMSE superior or equal to 20 were included. The included patients also presented chronic pain, assessed with Simple Verbal Scale27 at moderate or severe stage (score superior or equal to 2/4 since more than 1 month) and were able to complete the clinical and neuropsychological evaluations. Psychotropic drugs, acetylcholinesterase inhibitors, or memantine had to be stabilized for more than 3 months, and pain medication for at least 1 month.

Ethics Approval

The study protocol was reviewed and approved by Saint Etienne University Ethics Committee (Comité de Protection des Personnes). This committee encompasses the ethical approval of all 3 sites of our data collection. All procedures were in accordance with the Declaration of Helsinki and the International Conference on Harmonization Good Clinical Practice Guidelines. All participants signed an informed consent.

Measures

We considered the type of group with the intervention modalities being painting and singing as well as the time of the evaluation, that is, before or after each session.

Outcomes

1. The differences in immediate changes before and after in well-being as measured by EVIBE (in French: Evaluation Instantanée du Bien-Etre) and pain as measured by Numeric Rating Scale (NRS) between SI group sessions and PI group sessions.
2. The immediate change in well-being and pain before and after SI and PI sessions in SI and PI groups.

Outcome Measures

The EVIBE scale and the NRS have been chosen because these scales were sufficiently sensitive and adapted to patients with cognitive impairment.

The NRS28 was used as a measure of pain intensity at a given moment. It requires a lower degree of abstraction than the visual analog scale and can therefore be used in patients with early AD.29 Participants had to select the number that indicates the intensity of the pain felt at the moment. The
score varies from 0 (no pain) to 10 (the most severe pain one can imagine).

For the assessment of the immediate perceived well-being in patients with cognitive impairment, we used the EVIBE scale (in French: Evaluation Instantanée du Bien-Etre), a Visual Analog Scale. On a graduated ruler, the patients had to answer the question “How do you feel now?” The response varies from 0 (feeling bad) to 5 (feeling very good). To help the patient understand the instructions, pictograms representing simple facial expressions (joy, neutrality, and sadness) were proposed. The EVIBE scale can be used at severe stages of dementia and has already been used for measuring changes following a nondrug intervention. Furthermore, well-being measured by this scale was shown to be highly correlated with QoL.

Covariates
Sociodemographic and clinical assessment variables, including age, gender, and educational level, exploring global cognitive function, were collected at baseline and were used in the present analysis as covariates.

Randomization
All participants provided written inform consent. To ensure equal distribution between treatment groups, the randomization was made by a team member who had no contact with the patients. In each center (Lyon, Saint Etienne, see above), consented participants were randomly assigned to an SI or a PI group with a 1:1 ratio.

Blinding
All outcome measures were assessed by an independent psychologist in each center, who was different from the practitioner conducting SI or PI. The database and the statistical analysis were made by research team members. Both psychologists and the research team members were blind to the intervention type.

Intervention Conditions
The patients participated in 12 weekly 2-hour sessions during 3 months. They came to the hospital only for LACMé research. They had no other activities or contacts with other staff members.

Singing intervention groups. The SI was delivered by a professional choir conductor accompanied by a psychologist, whose role was to help the conductor to manage the patients if necessary (in particular welcome, departure, and possible difficulties with patients during the sessions). Before the intervention, the patients assigned to SI groups were asked to fulfill a questionnaire collecting their musical preferences among several songs. The choir conductor selected the songs according to the patients’ preferences. After a personalized welcome, the patients performed a body and voice warm-up before song learning. Four different songs previously chosen by the patients among a list of well-known songs were practiced across the different sessions. The songs were then worked by the patients with piano accompaniment made by the choir conductor.

Painting intervention groups. The PI was done by a painting teacher accompanied by a psychologist. After a personalized welcome in the first part of each session, paintings of professional painters were shown to the patients, generating discussions across the group. The second part of the session was devoted to the patients’ painting realization according to a predetermined theme.

Data Collection
The pain and well-being levels were collected by an independent psychologist before and after each session. The duration for these evaluations was approximately 5 minutes.

Data Analysis
The mean value of EVIBE and NRS before and then after the sessions was computed for each participant. The evolution of EVIBE and NRS scores across the sessions was then compared between SI and PI groups using mixed-effects models, including time as a fixed effect and participants as a random effect. The evolution of NRS and EVIBE was then compared across each session, between the 2 groups, with mixed-effects models. All statistical tests were 2 tailed. A P values inferior to .05 was considered as significant. Statistical analyses were performed with SPSS version 17 (SPSS Software, Chicago, Illinois).

### Table 1. Patients’ Baseline Characteristics.a

| Patients’ Characteristics | Singing Intervention (N = 31) n (%) | Painting Intervention (N = 28) n (%) |
|---------------------------|-----------------------------------|-----------------------------------|
| Gender                    |                                    |                                    |
| Women                     | 23 (74.2)                          | 16 (57.1)                          |
| Education                 |                                    |                                    |
| <5 years                  | 2 (6.5)                            | 5 (17.9)                           |
| 5-6 years                 | 8 (25.8)                           | 8 (28.6)                           |
| 7-8 years                 | 10 (32.3)                          | 6 (21.4)                           |
| 9-11 years                | 5 (16.1)                           | 2 (7.1)                            |
| >11 years                 | 6 (19.4)                           | 7 (25.0)                           |
| Mean (SE)                 |                                    |                                    |
| Age                       | 78.8 (7.43)                        | 80.2 (5.71)                        |
| MMSE                      | 25.07 (2.26)                       | 24.18 (2.72)                       |
| SVS                       | 2.06 (0.24)                        | 2.17 (0.38)                        |

Abbreviations: MMSE, Mini-Mental State Examination; SVS, Simple Verbal Scale. 
*aSVS varying from 0 to 4, with 0 indicating no pain, and 4 indicating very intense pain.
Results

Population’s Characteristics

A total of 59 patients completed the baseline evaluation. Among them, 31 received SI and 28 PI. The demographic and clinical characteristics of the samples at baseline are summarized in Table 1. There was no imbalance in variables between groups with regard to demographic and severity of cognitive disorder.

Pain and Well-Being Assessment

Table 2 shows the results of pain and well-being measures including means and standard deviations for NRS and EVIBE scales before and after sessions, and the results of the mixed models adjusted on age, sex, and educational level. The NRS scores decreased over time; however, the results did not reach the level of significance. No difference between groups or interaction time by group was observed in these analyses. The mean EVIBE score increased over time in both SI and PI. A trend toward significance was observed for the group factor. However, there was no interaction between time and group in this analysis.

Well-Being Assessment for Each Session

Figure 1 shows more precisely the evolution of EVIBE scores over time for each session in both SI and PI. The mean EVIBE score increased over time across each session in both SI and PI. The increase was significant for 9 sessions, and a trend toward increasing was observed in the 3 remaining sessions. There is not group effect except for session 4, for which there is a significant difference between groups. This analysis shows no interaction between time and group.

Discussion

This report is one of the few to assess the immediate benefit of artistic interventions over several sessions with a randomized trial. The findings reveal an improvement in well-being for both SI and PI groups after each session, compared to the
assessment before the session. However, we did not observe this improvement across the sessions for pain intensity measurement. As in the previous study, we did not observe a stronger benefit of SI compared to PI for pain and well-being.

Our present findings confirm the usefulness of the EVIBE scale, allowing to reveal the patients’ well-being in the present moment. Moreover, this scale seems to be sensitive to change and to show a benefit on well-being after a nonpharmacological care activity. This scale is thus accessible to patients with cognitive disorders and has the advantage of being easy and fast to use.

The results of the EVIBE scale revealed a general benefit of artistic activity on patients’ well-being, regardless of the type of artistic practice. Similarly, previous researches have shown that artistic activities, whether musical or visual, can improve not only behavior and increase the sociability of patients with dementia but also well-being and QoL. An improvement of about 0.5 points on the EVIBE scale is observed after the activities which seems to be clinically meaningful. However, as the EVIBE scale is recent, no relevance threshold has been determined.

When considering the evolution of pain judgments across the sessions, we can see a slight to moderate improvement, but the results were not significant. The painful intensity at the present moment measured with NRS scale may not be relevant for assessing change in painful feeling. Indeed, previous studies have shown that ratings of pain intensity remain often stable over time. A patient might judge pain with a relatively constant score even though an overall improvement is felt.

Otherwise, our previous results revealed an improvement in several measures including both usual and intense pain and pain repercussions. These results were revealed after the 12 sessions compared to the baseline both for SI and PI. Although a significant immediate benefit on pain could not be shown here, it nevertheless seems to be obtained in a longer term assessment.

Some assertions can explain the differences with our initial conclusions. Indeed, after the 12 sessions, we used a numerical scale and required participants to base their judgment on the maximum intensity of the last 7 days and not at the present time, as done in this study. The way the question is asked implies that the patient appeals more to his/her memory of pain than to the currently felt pain. This is also true for the other measures used after the 12 intervention sessions: the pain repercussions and the usual pain on the last 7 days. Memory of pain has been shown to be strongly linked to the emotional context and consequently to the feeling of well-being. The results obtained after a long time period from the sessions are therefore difficult to dissociate from a more general improvement in well-being. Even if the interventions are more beneficial on the memory of pain than on the present pain per se, these results remain interesting as studies showed that this memory of pain can influence the chronicity of the pain afterward.

The present findings must be interpreted within the context of several limitations. First and foremostly, the nonsignificant results on the pain scale are probably due in part to the variability in pain scores between patients, as evidenced by the wide standard deviations observed in our results. Variability is also observed between baseline scores and the pain scores measured during the sessions, where some patients reported low pain levels. This may have limited statistical power over pain intensity measures. In the future, it would be interesting to propose a similar study with patients reporting higher pain scores but nevertheless comparable etiologies.

In addition, we did not include a control group with usual care or nonartistic activity. This kind of control group would have been interesting to support the hypothesis that artistic activities have a direct benefit on well-being rather than other activities. However, we chose to propose a painting group as a comparison group because the main objective of our work was to highlight the specificity of singing compared to another artistic activity. Indeed, unlike painting, a robust scientific literature shows the interest of music on different emotional and physiological parameters. We expected more profit with SI than with other art activities. This exploratory study shows an improvement in well-being but does not allow differentiating these results from a single placebo or a social effect. Moreover, the randomized controlled trial model is scientifically relevant but does not allow to adapt the groups according to patient’s preferences, which is usually done in routine care. Even though the interventions took into account their preferences they could have been more effective if the patients had been able to choose between SI and PI. Additional data are essential to determine whether the benefit is related to the artistic nature of these activities.

In conclusion, these exploratory results suggest that PI and SI provide an immediate benefit on the patients’ well-being. Immediate benefit is often sought in the management of dementia, as for example, in nursing homes or day care centers. These results encourage art therapy initiatives as a nonpharmacological method to instantly improve the well-being of patients with cognitive impairment.

Authors’ Note
Pierre Krolak-Salmon is also affiliated with University Lyon, Lyon, France.

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ORCID iD
Elodie Pongan https://orcid.org/0000-0001-8674-6361

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