Mannose 6-phosphonate labelling: A key for processing the therapeutic enzyme in Pompe disease
Anastasia Godefroy, Morgane Daurat, Afitz da Silva, Ilaria Basile, Khaled El Cheikh, Catherine Caillaud, Sabrina Sacconi, Benedikt Schoser, Henry-vincent Charbonné, Magali Gary-bobo, et al.

To cite this version:
Anastasia Godefroy, Morgane Daurat, Afitz da Silva, Ilaria Basile, Khaled El Cheikh, et al.. Mannose 6-phosphonate labelling: A key for processing the therapeutic enzyme in Pompe disease. Journal of Cellular and Molecular Medicine, Wiley Open Access, 2019, 23 (9), pp.6499 - 6503. 10.1111/jcmm.14516 . hal-03116444

HAL Id: hal-03116444
https://hal.archives-ouvertes.fr/hal-03116444
Submitted on 20 Jan 2021

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
SHORT COMMUNICATION

Mannose 6-phosphonate labelling: A key for processing the therapeutic enzyme in Pompe disease

Anastasia Godefroy1,2 | Morgane Daurat1,2 | Afitz Da Silva1,2 | Ilaria Basile2 | Khaled El Cheikh2 | Catherine Caillaud3 | Sabrina Sacconi4 | Benedikt Schoser5 | Henry-Vincent Charbonné2 | Magali Gary-Bobo1 | Alain Morère1 | Marcel Garcia1,2 | Marie Maynadier2

1IBMM, CNRS, ENSCM, University of Montpellier, Montpellier, France
2NanoMedSyn, Montpellier, France
3Biochimie Météabolique et Protéique, AH-HP, Hôpital Necker Enfants-Malades and Inserm U1151, Institut Necker Enfants Malades, Université Paris-Descartes, Paris, France
4Service Système Nerveux Périphérique, Muscle et SLA, Centre Hospitalier Universitaire de Nice, Centre National de la Recherche Scientifique, Institut National de la Santé et de la Recherche Médicale, Institute for Research on Cancer and Aging of Nice, Université Côte d’Azur, Nice, France
5Department of Neurology, Friedrich-Baur-Institute, Ludwig-Maximilians University Munich, Munich, Germany

Abstract
In the search of a better enzyme therapy in Pompe disease, the conjugation of mannose 6-phosphonates to the recombinant enzyme appeared as an enhancer of its efficacy. Here, we demonstrated that the increased efficacy of the conjugated enzyme is partly due to a higher intracellular maturation because of its insensitivity to acid phosphatases during the routing to lysosomes.

KEYWORDS
acid phosphatases, acid α-glucosidase, enzyme replacement therapy, intracellular processing, lysosomal storage disease, mannose 6-phosphate receptor

This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.

© 2019 The Authors. Journal of Cellular and Molecular Medicine published by John Wiley & Sons Ltd and Foundation for Cellular and Molecular Medicine.
1 | INTRODUCTION

In Pompe disease, the current enzyme replacement therapy based on a human recombinant acid α-glucosidase (rhGAA) naturally bearing mannose 6-phosphate (M6P) has a limited efficacy.1–4 We recently proposed a conjugation of rhGAA with a phosphonate analogue of mannose 6-phosphate (called AMFA) which enables never-before seen improvements on walking ability and on musculoskeletal health in the aged Pompe mouse model.5,6 Here, we hypothesized that this new therapeutic efficacy is not solely due to a better internalization via M6P receptor pathway and we investigated the enzyme processing. The intracellular maturation of rhGAA (110 kD) is complex and consists in successive proteolyses up to an endosomal 95 kD intermediate form, a 76 kD active form and a 60‐70 kD lysosomal mature form (Scheme S1).7,8 The 76 and 60‐70 kD GAA species show a 7‐10‐fold increased activity.7 Bali et al.9 clearly established on patient biopsies that the overall activity was correlated with the presence of the 60‐70 kD form.

Secondly, since an overexpression of acid phosphatases has been observed in several lysosomal storage disorders (LSD)9,10 we also analysed the involvement of lysosomal acid phosphatases ACP2 and ACP5 in rhGAA maturation.

2 | MATERIALS AND METHODS

See Supporting Information for detailed description.

3 | RESULTS

3.1 | AMFA grafting on rhGAA enables in vitro and in vivo complete enzyme processing

A high uptake of 110 kD GAA precursor was observed in cultured myoblasts from adult Pompe patients for both rhGAA and rhGAA-AMFA treatments (Figure 1A); however, 76 kD form (Figure S1A) and 60‐70 kD form (Figure S1B) are 3‐ and 5‐fold more important for rhGAA-AMFA treated cells. As expected intracellular GAA activity was also significantly increased for rhGAA-AMFA as compared to rhGAA (Figure S1C). Similar results were found on different myoblasts (Figure S1D–G). GAA genetic mutations of the corresponding primary cultured cells are listed in Table S1.

We then differentiated myoblasts into myotubes according to the technique described by Nascimbeni et al 2012.11 In myotubes, the 76 kD and 60‐70 kD forms were significantly increased by rhGAA-AMFA treatment (4.1‐ and 2.2‐fold) as compared to rhGAA (Figure 1B‐C).

We also analysed the enzyme maturation in 10‐month‐old Pompe mice treated weekly with 5 mg/kg rhGAA, rhGAA-AMFA or vehicle during 3 months. While no specific band was detected in the control Pompe tissue, the 95 kD intermediary form was detected after rhGAA treatment (Figure 1D). In mice injected with rhGAA-AMFA, the 95 kD form, the 76 kD active form and/or the 60‐70 kD mature form were observed. This higher maturation was also associated with a gain in GAA activity in muscle biopsies (data not shown). Together, these results indicate that AMFA conjugation increases rhGAA maturation both in patient cultured cells and in aged Pompe mice.

FIGURE 1 | Uptake and maturation of rhGAA and rhGAA-AMFA in myoblasts and myotubes of Pompe patients (P12, P14) and in quadriceps of treated Pompe mice. A, P12 myoblasts. B, C, myotubes differentiated from myoblasts of adult (P12) and juvenile (P14) patients were incubated with 50 nmol/L rhGAA, rhGAA-AMFA or with vehicle (Control) for 8 h (myoblasts) or 48 h (myotubes) in culture medium. The cell extracts (5 µg) were analysed by Western blot using human GAA or actin antibodies. D, Maturation of rhGAA and rhGAA-AMFA in quadriceps of aged Pompe mice. The tests were performed on 10‐month‐old mice treated with 5 mg/kg/week of rhGAA or rhGAA-AMFA or by vehicle (Control), during 13 weeks. The maturation of the enzymes in quadriceps on aged Pompe mice is studied by Western blot on 20 µg tissue extract using human GAA or actin antibodies. Black arrows indicate respectively 110 kD (inactive precursor), 95 kD (inactive intermediary), 76 kD (active intermediary) and 60‐70 kD GAA (mature active) forms and actin is a control for total protein loading.
3.2 Impact of acid phosphatases in GAA processing

The main difference between AMFA and Mannose 6-Phosphate is the replacement of the phosphate moiety by a phosphonate group insensitive to phosphatase in contrast to M6P. In agreement with a previous study, we found that acid phosphatases activity is significantly increased in Pompe patient myoblasts (264 ± 96%) as compared to normal myoblasts (value set as 100 ± 11%, P < 0.005 Student’s t test) (Figure 2A). As such phosphatases could be involved
in M6P signal deterioration on rhGAA, we demonstrated that the inhibition of overexpressed phosphatases with sodium fluoride or \(\beta\)-glycerophosphate allowed the processing of rhGAA into 76 kD and 60-70 kD active forms (Figure 3).

Expressions of ACP2 and ACP5, two major acid phosphatases involved in M6P dephosphorylation\(^7\) were assessed (Figure 2B). ACP2 is overexpressed in almost all Pompe disease samples, whereas few ones overexpressed ACP5.

The inhibition with specific siRNAs of 58% ACP2 and 24% ACP5 expression (Figure 2C-E) was sufficient to enhance rhGAA maturation (Figure 2D-F) and partially that of the endogenous deficient enzyme.

## DISCUSSION

In this report, significant differences in the processing of rhGAA-AMFA and rhGAA are presented. Although both enzymes are well internalized, only rhGAA-AMFA undergoes cleavage to 76 kD active and 60-70 kD mature forms in primary cultures of fibroblasts, myoblasts and myotubes from Pompe patients and in aged Pompe mice after 3-month treatment. These data establish that the protease machinery necessary for enzymatic maturation is still functional in both adult patients and aged Pompe mice which were previously considered as ERT refractory.\(^4\)

Then, we considered the role of phosphatases in GAA maturation. In LSD, several reports evidence an overexpression of lysosomal acid phosphatases. As already demonstrated for lysosomal enzymes in cancers,\(^12,13\) we supposed that overexpressed phosphatases can overflow from lysosomes to endo-lysosomal vesicles. Such abnormally localized phosphatases could prevent GAA-M6PR complexing before the enzyme reaches the lysosomes and thus impair enzyme endo-lysosomal maturation. Our present analysis indicates an increase of the acid phosphatases activity and more specifically, an elevation of ACP2 and ACP5. Using general inhibitors and specific siRNAs, we obtained a partial phosphatases inhibition sufficient to allow the formation of rhGAA active and mature forms. Although different cell types remain to be evaluated, these data already evidence that acid phosphatases overexpression prevents rhGAA processing.

The outcome of the unmatured rhGAA was not investigated here. However, aberrant localization of rhGAA into autophagosomes could be hypothesized.\(^14\)

In conclusion, the higher therapeutic efficacy of rhGAA-AMFA observed in vitro and in vivo\(^5,6\) is associated with an increase of enzyme maturation. Altogether, these data suggest that AMFA targeting may represent a potential therapeutic advantage for Pompe disease and also for other LSD which overexpress acid phosphatases.

## ACKNOWLEDGEMENTS

We thank Dr Gilles CARNAC, from Inserm U1046-UMR CNRS 9214, and Pr François RIVIER, from the CHU Montpellier-Guy de Chauliac, for their scientific and medical advices. This work was supported by French National Research Agency ANR-13-RPIB-0012, by Association Nationale Recherche Technologie grant 2016/0628 and grant 2016/0629, by APPI grant N°DOS0026362/00 from BPI, by NanoMedSyn and by Occitanie-Midi Pyrénées Region grant PILE 180060 and PILE 180064, by Vaincre les Maladies Lysosomales Suisses. For sample providing, we thank Pr Schaeffer from CBC Biotec Biobank, the Reference Center for Neuromuscular Diseases and ALS of Nice University Hospital, and the Muscle Tissue Culture Collection MTCC. We thank the Biobank of Cells, Tissues and DNA from patients with neuromuscular diseases, member of the Telethon Network of Genetic Biobanks (project no. GTB12001), funded by Telethon Italy, and of the EuroBioBank network, for providing us specimens. The Muscle Tissue Culture Collection is part of the German network on muscular dystrophies (MD-NET, service structure S1, 01GM0601) and the German network for mitochondrial disorders (mito-NET, project D2, 01GM0862) funded by the German Ministry of Education and Research (BMBF, Bonn, Germany). The Muscle Tissue Culture Collection is a partner of EuroBioBank (www.eurobiobank.org) and TREAT-NMD (www.treat-nmd.eu). AG, MD, ADS are CIFRE PhD students from Association Nationale Recherche Technologie. IB, KEC, HVC, MG, MM are employees of NanoMedSyn. MG, MGB and AM are co-founders and have received consulting fees from NanoMedSyn.

## CONFLICT OF INTEREST

The authors declare that they have no conflicts of interest with the contents of this article.

## DATA AVAILABILITY

The data that support the findings of this study are available from the corresponding author upon reasonable request.

## ORCID

Alain Morère \(\text{ID} \) https://orcid.org/0000-0002-3269-5172

## REFERENCES

1. van der Ploeg AT, Clemens PR, Corzo D, et al. A Randomized study of alglucosidase alfa in late-onset Pompe’s disease. *N Engl J Med*. 2010;362:1396-1406.
2. Schoser B, Stewart A, Kanters S, et al. Survival and long-term outcomes in late-onset Pompe disease following alglucosidase alfa treatment: a systematic review and meta-analysis. *J Neurol*. 2017;264:621-630.
3. Maga JA, Zhou J, Kambampati R, et al. Glycosylation-independent lysosomal targeting of acid \(\alpha\)-glucosidase enhances muscle glycogen clearance in Pompe mice. *J Biol Chem*. 2013:288:1428-1438.
4. Zhu Y, Jiang J-L, Gumlaw NK, et al. Glycoengineered acid \(\alpha\)-glucosidase with improved efficacy at correcting the metabolic aberrations and motor function deficits in a mouse model of Pompe disease. *Mol Ther*. 2009;17:954-963.
5. El Cheikh K, Basile I, Da Silva A, et al. Design of potent mannose 6-phosphate analogues for the functionalization of lysosomal enzymes to improve the treatment of Pompe disease. Angew Chem Int Ed Engl. 2016;55:14774-14777.
6. Basile I, Da Silva A, El Cheikh K, et al. Efficient therapy for refractory Pompe disease by mannose 6-phosphate analogue grafting on acid α-glucosidase. J Control Release. 2018;269:15-23.
7. Moreland RJ, Jin X, Zhang XK, et al. Lysosomal acid α-glucosidase consists of four different peptides processed from a single chain precursor. J Biol Chem. 2005;280:6780-6791.
8. Bali DS, Tolun AA, Goldstein JL, Dai J, Kishnani PS. Molecular analysis and protein processing in late-onset Pompe disease patients with low levels of acid α-glucosidase activity. Muscle Nerve. 2011;43:665-670.
9. Tsuburaya RS, Monna K, Oya Y, et al. Acid phosphatase-positive globular inclusions is a good diagnostic marker for two patients with adult-onset Pompe disease lacking disease specific pathology. Neuromuscul Disord. 2012;22:389-393.
10. Makrypidi G, Damme M, Muller-Loennies S, et al. Mannose 6 dephosphorylation of lysosomal proteins mediated by acid phosphatases Acp2 and Acp5. Mol Cell Biol. 2012;32:774-782.
11. Nascimbeni AC, Fanin M, Masiero E, Angelini C, Sandri M. The role of autophagy in the pathogenesis of glycogen storage disease type II (GSDII). Cell Death Differ. 2012;19:1698-1708.
12. Rochefort H, Liaudet E, Garcia M. Alterations and role of human cathepsin D in cancer metastasis. Enzyme Protein. 1996;49:106-116.
13. Aggarwal N, Sloane BF. Cathepsin B: multiple roles in cancer. Proteom Clin Appl. 2014;8:427-437.
14. Raben N, Ralston E, Chien Y-H, et al. Differences in the predominance of lysosomal and autophagic pathologies between infants and adults with Pompe disease: implications for therapy. Mol Genet Metab. 2010;101:324-331.

SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.

How to cite this article: Godefroy A, Daurat M, Da Silva A, et al. Mannose 6-phosphonate labelling: A key for processing the therapeutic enzyme in Pompe disease. J Cell Mol Med. 2019;23:6499–6503. https://doi.org/10.1111/jcmm.14516