Pathophysiology and targets for treatment in hereditary galactosemia: A systematic review of animal and cellular models

Minela Haskovic1,2,3 | Ana I. Coelho1,2,3 | Jörgen Bierau2 | Jo M. Vanoevelen2,3 | Laura K. M. Steinbusch2 | Luc J. I. Zimmermann1,3 | Eduardo Villamor-Martinez1,3 | Gerard T. Berry4 | M. Estela Rubio-Gozalbo1,2,3

1Department of Pediatrics, Maastricht University Medical Center+, Maastricht, The Netherlands
2Department of Clinical Genetics, Maastricht University Medical Center+, Maastricht, The Netherlands
3GROW-School for Oncology and Developmental Biology, Maastricht University, Maastricht, The Netherlands
4The Manton Center for Orphan Disease Research, Division of Genetics and Genomics, Boston Children’s Hospital, Harvard Medical School, Boston, Massachusetts

Correspondence
M. E. Rubio-Gozalbo, Department of Pediatrics, Maastricht University Medical Center+, P. Debyelaan 25, P.O. Box 5800, 6202 AZ Maastricht, The Netherlands. Email: estela.rubio@mumc.nl

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Abstract
Since the first description of galactosemia in 1908 and despite decades of research, the pathophysiology is complex and not yet fully elucidated. Galactosemia is an inborn error of carbohydrate metabolism caused by deficient activity of any of the galactose metabolising enzymes. The current standard of care, a galactose-restricted diet, fails to prevent long-term complications. Studies in cellular and animal models in the past decades have led to an enormous progress and advancement of knowledge. Summarising current evidence in the pathophysiology underlying hereditary galactosemia may contribute to the identification of treatment targets for alternative therapies that may successfully prevent long-term complications. A systematic review of cellular and animal studies reporting on disease complications (clinical signs and/or biochemical findings) and/or treatment targets in hereditary galactosemia was performed. PubMed/MEDLINE, EMBASE, and Web of Science were searched, 46 original articles were included. Results revealed that Gal-1-P is not the sole pathophysiological agent responsible for the phenotype observed in galactosemia. Other currently described contributing factors include accumulation of galactose metabolites, uridine diphosphate (UDP)-hexose alterations and subsequent impaired glycosylation, endoplasmic reticulum (ER) stress, altered signalling pathways, and oxidative stress. galactokinase (GALK) inhibitors, UDP-glucose pyrophosphorylase, and unfolded protein response.

Abbreviation: ATP, adenosine triphosphate; BiP, binding protein; CG, classic galactosemia; CHO, Chinese hamster ovary; ER, endoplasmic reticulum; Gal-1-P, galactose-1-phosphate; GALE, UDP-galactose 4’-epimerase; GALK, galactokinase; GALM, galactose mutarotase; GALT, galactose-1-phosphate uridylyltransferase; Glc-1-P, glucose-1-phosphate; HPLC, high-performance liquid chromatography; PI3/Akt, phosphatidylinositol-4,5-bisphosphate 3-kinase/protein kinase B; PRISMA, Preferred Reporting Items for Systematic Reviews and Meta-Analysis; SIGN, Scottish Intercollegiate Guidelines Network; SNA, Sambucus nigra agglutinin; SOD, superoxide dismutase; UDP-Gal, uridine diphosphate-galactose; UDP-GalNAc, UDP-N-acetylgalactosamine; UDP-Glc, uridine diphosphate-glucose; UDP-GlcNAc, UDP-N-acetylglucosamine; UGP, UDP-glucose pyrophosphorylase; UPR, unfolded protein response.

Minela Haskovic and A. I. Coelho are co-first authors.
G.T. Berry and M. E. Rubio-Gozalbo are co-last authors.

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Pyrophosphorylase (UGP) up-regulation, uridine supplementation, ER stress reducers, antioxidants and pharmacological chaperones have been studied, showing rescue of biochemical and/or clinical symptoms in galactosemia. Promising co-adjuvant therapies include antioxidant therapy and UGP up-regulation. This systematic review provides an overview of the scattered information resulting from animal and cellular studies performed in the past decades, summarising the complex pathophysiological mechanisms underlying hereditary galactosemia and providing insights on potential treatment targets.

KEYWORDS
animal models, cellular models, hereditary galactosemia, pathophysiology, treatment targets

1 | INTRODUCTION

Despite that the first description of galactosemia that dates from more than 100 years ago (1908) and decades of research, the pathophysiology of this disease is complex and not yet fully understood. Studies in cellular and animal models in the past decades have led to an enormous progress and advancement of knowledge. Systematically summarising current evidence in the pathophysiology underlying hereditary galactosemia may contribute to the identification of treatment targets for alternative therapies that may successfully prevent the occurrence of long-term complications.

Galactose is an aldohexose and is mainly metabolised via the Leloir pathway (Figure 1). In most organisms, the first step of the Leloir pathway involves epimerization of β-D-galactose to α-D-galactose by galactose mutarotase (GALM; EC 5.1.3.3), encoded by the GALM gene located on chromosome 2p22.1. The next step entails the phosphorylation of α-D-galactose into galactose-1-phosphate (Gal-1-P) by the action of galactokinase (GALK; EC 2.7.1.6), encoded by the GALK1 gene located on chromosome 17q25.1. Together with uridine diphosphate-glucose (UDP-Glc), Gal-1-P is converted to glucose-1-phosphate (Glc-1-P) and uridine diphosphate-galactose (UDP-Gal) by galactose-1-phosphate uridylyltransferase (GALT; EC 2.7.7.12) encoded by the GALT gene located on chromosome 9p13. UDP-galactose 4’-epimerase (GALE; EC 5.1.3.2) is encoded by the GALE gene located on chromosome 1p36.11 and catalyses the conversion of UDP-Gal into UDP-Glc. In mammals, GALE is also responsible for the interconversion of UDP-N-acetylglucosamine (UDP-GlcNAc) and UDP-N-acetylgalactosamine (UDP-GalNAc). GALE maintains the intracellular pools of UDP-Gal/UDP-Glc and UDP-GlcNAc/UDP-GalNAc.

Galactosemia is an inborn error of carbohydrate metabolism caused by a severe impairment of any of the enzymes involved in the Leloir pathway and comprises four subtypes. Type 1, also known as classic galactosemia (CG, OMIM 230400) is characterised by severe deficiency of the GALT enzyme. Glycosylation abnormalities have been described in CG, characterised by processing defects (reviewed by Maratha et al2). UDP-hexoses abnormalities and inhibition of galactosyltransferases by high levels of Gal-1-P have been suggested as potential mechanisms.3-6 Defective galactosylation has been described in patient fibroblasts7,8 and serum glycoproteins of untreated CG patients.3,9 CG is potentially lethal in the neonatal period, with multi-organ involvement. The current standard of care, a galactose restricted diet, although life-saving in the neonatal period, fails to prevent complications, such as cognitive impairments, neurological symptoms, and subfertility in female patients.10 Type 2 is characterised by deficiency of the GALK enzyme (OMIM 230200), in untreated patients presenting with cataract.11 Other symptoms have also been described in several patients: pseudotumor cerebri, hypoglycemia (asymptomatic), mental retardation, microcephaly, failure to thrive, seizures, deafness, hepatomegaly, and hypercholesterolemia.12,13 Type 3 is caused by a deficiency of the GALE enzyme (OMIM 230350), presenting with acute symptoms similar to CG (14 updated in 2016). Long-term outcomes reported in GALE deficient patients include physical and cognitive developmental delay and/or learning disabilities. Recently, the fourth type of galactosemia has been described, caused by biallelic pathogenic variants in GAL.15

When one of the Leloir pathway enzymes is deficient, galactose is disposed through alternative pathways. These include galactose reduction to galactitol, catalysed by the NADPH dependent aldose reductase (AR, EC 1.1.1.21), galactose oxidation, leading to the production of galactonate by galactose dehydrogenase (EC 1.1.1.48) and presumably activation of the pyrophosphorylase pathway, which converts galactose to UDP-Glc by the sequential activities of GALK and UDP-glucose pyrophosphorylase (UGP, EC 2.7.7.10).16
In the past decades, a myriad of research has taken place using cellular and animal models to gain insight on the complex pathophysiology of hereditary galactosemia. This study provides a comprehensive systematic review of the scattered information in the different studies, through long periods of time. This can be of great benefit to researchers in the field and provide insights on potential treatment targets.

2 | MATERIALS AND METHODS

This study was carried out and reported according to the Preferred Reporting Items for Systematic Reviews and Meta-Analysis (PRISMA) guidelines. The PRISMA checklist for this report can be found in the Supplementary file 1. A protocol was developed prospectively that detailed the specific objectives, criteria for study selection and the approach to assess study quality and outcomes. The protocol was first published on October 16th, 2018. During the conduction of this systematic review, the protocol was adapted and updated on August 16th, 2019 (http://syrf.org.uk/protocols/).

2.1 | Research question

The primary outcomes for this systematic review were disease complications in GALK, GALT, and/or GALE deficiency studied in animal and/or cellular models of the disease, reported as clinical signs and/or biochemical findings. Our aim was to answer the following question: “What have cellular and/or animal models contributed to unravel the pathophysiology of hereditary galactosemia and to identify targets/approaches for treatment?”
2.2 | Search strategy

A computerised literature search was conducted in PubMed/MEDLINE, EMBASE, and Web of Science by two independent researchers (MH and AIC), using animal filters.¹⁸,¹⁹ Databases were searched on October 24th, 2018. Additionally, reference lists from included studies were hand-searched to identify relevant studies. The records retrieved were imported and de-duplicated in ENDNOTE X8. The complete search strategies can be extracted from the (Supplementary file 2).

2.3 | Inclusion and exclusion criteria for study selection

All original studies including animal and/or cellular models focusing on disease complications and/or treatment targets in hereditary galactosemia were included. Articles in a language other than English, case reports, conference abstracts, and reviews were excluded.

2.4 | Study eligibility

Titles and abstracts were independently screened by M.H. and A.I.C. Studies not relating to the research question were excluded. Full-text articles were independently reviewed by M.H. and A.I.C. for the inclusion in the systematic review. In case of exclusion, the reason was reported. In case of a discrepancy in the selection of studies, considerations were discussed with the other authors until consensus on eligibility was reached.

2.5 | Data collection

Data were independently extracted by M.H. and A.I.C. according to predefined criteria in the protocol. Outcome measures of interest were disease complications, reported as clinical signs (mimicking the human phenotype: neurologic and cognitive impairment [motor abnormalities], gonadal damage [reduced fertility], cataract, as well as survival rate) and/or biochemical findings (increased metabolites [galactose, Gal-1-P, galactitol, galactonate], decreased enzyme activity [GALT, GALK, GALE], altered levels of inositol [and related metabolites], disrupted signalling pathways, glycosylation abnormalities/altered UDP-hexoses levels, oxidative stress, activation of the unfolded protein response [UPR], endoplasmic reticulum [ER] stress, apoptosis).

2.6 | Risk of bias and quality assessment in individual studies

Two independent researchers (M.H. and A.I.C.) performed and discussed risk of bias assessment using the SYRCLE’s risk of bias tool for animal studies²⁰ and the Scottish Intercollegiate Guidelines Network (SIGN, https://www.sign.ac.uk/checklists-and-notes.html) quality appraisal checklists for case-control study designs for cellular studies. Articles were appraised as low, acceptable, or high quality. Those assessed as low quality were excluded from the review. Discrepancies were resolved by discussion with all authors.

3 | RESULTS AND DISCUSSION

3.1 | Included studies

The computerised literature search identified 3407 potential hits, from which 46 articles were included in this systematic review. Figure 2 shows the PRISMA flow diagram for inclusion of individual studies. Of the 46 included articles, three reported on GALK deficiency, 39 on GALT deficiency, and 10 on GALE deficiency. Synoptic tables of the included studies are presented in the supplementary material (Supplementary file 3).

3.2 | Quality assessment

The quality of the included individual studies was assessed using SYRCLE’s risk of bias tool and SIGN quality appraisal checklist (Supplementary file 4).

We categorised the obtained data on pathophysiology (summary in Tables 1 and 2, and Figure 1) and treatment targets (summary in Table 3 and Figure 1) as follows: (a) accumulation of metabolites, (b) UDP-hexoses abnormalities and glycosylation impairments, (c) signalling pathways alterations, UPR activation, ER stress, and oxidative stress, (d) structural and functional defects, and (e) treatment approaches.

In the past decades, studies in patient cell lines, yeast—Saccharomyces cerevisiae—²¹ bacteria—Escherichia coli—²²-²⁴ as well as studies in animal models have been performed. The first animal model of classic galactosemia, a Galt knockout mouse, was described in 1996.²⁵ After several years, a mouse model for GALK deficiency was developed.²⁶ Fruit fly—Drosophila melanogaster—models for both GALT and GALE deficiency followed in 2010.²⁷,²⁸ Furthermore, a nematode—Caenorhabditis elegans—for GALE
deficiency was developed. In addition to the mouse model described by Leslie et al., a Galt gene-trapped mouse was reported in 2014. More recently, a zebrafish—Danio rerio—model for GALT deficiency was described.

3.3 Accumulation of metabolites

3.3.1 GALK deficiency

In GALK deficiency, there is no accumulation of Gal-1-P. Ross et al. showed that GALK-deficient yeast demonstrated normal growth, no accumulation of Gal-1-P and normal UDP-Gal and UDP-Glc levels. Studies in fruit flies demonstrated that, despite low levels of Gal-1-P, loss of GALK slightly lowered survival rates, in both the absence and presence of dietary galactose. Loss of GALK had a larger negative impact on the climbing ability and fecundity, in the absence of galactose. Studies in a mouse model for GALK deficiency showed that increased galactitol synthesis, when overexpressing aldose reductase, led to cataract formation. A low expression of aldose reductase in the lens of mice has been reported. Moreover, galactitol accumulation depletes NADPH levels and leads to decreased glutathione reductase activity. As a result, free radicals accumulate, causing oxidative stress that can lead to cell death.

3.3.2 GALT deficiency

Gal-1-P has been hypothesized as one of the key pathogenic agents in classic galactosemia and studied extensively. Leslie’s GALT-deficient mouse demonstrated accumulated levels of Gal-1-P (in liver homogenates and erythrocytes) and galactose (in plasma) during the suckling period. Despite this metabolic abnormality, these mice did not exhibit the expected clinical phenotype, even after a high galactose diet. Substantial levels of Gal-1-P and galactitol in liver,肾 and brain of the adult mice were found and only a trace of galactitol in plasma. Highest levels of galactitol were reported in muscle and...
endogenous galactose production. Galactose level in the liver appeared to be lower than that in plasma, kidney and brain, likely due to its conversion to galactonate in the liver. Furthermore, a relatively low brain tissue galactose level was found which led to the hypothesis that the blood-brain barrier may restrict galactose entry. Galactonate has been measured in different tissues (liver, kidney, and brain) in mice. They observed that galactonate levels accumulated in all tissues after galactose exposure. In the unexposed situation, no galactonate was found in the brain. In another study, galactonate levels in mice muscle and heart were found 10 times higher than that in brain or kidney after galactose exposure. The combined accumulation of Gal-1-P and galactitol was suggested to cause the phenotype of human GALT deficiency.

In 2014, a Galt gene-trapped mouse was developed by Tang et al. They showed that galactose challenge in homozygous Galt gene-trapped lactating females affected the outcome of the newborn Galt gene-trapped mice. The newborn mice showed a high mortality rate, insults in brain tissues and manifestations of oxidative stress in red blood cells (altered glutathione/oxidised glutathione ratio). Brain lesions in the galactose-intoxicated pups were characterised by smaller, more acidophilic Purkinje cells and a thicker outer granular cell layer of the cerebellum. This finding led the authors to suggest a post-natal role of galactose in the pathophysiology of the ataxia observed in patients. The mice accumulated only modest levels of Gal-1-P in red blood cells, even under galactose challenge. Furthermore, they observed growth restriction in unexposed mice and reduced fertility in female mice who were not exposed to galactose beyond the weaning period. Authors proposed that this finding may explain the higher galactose resistance in rodents compared to humans.

Extensive studies in the GALT-deficient D. melanogaster revealed that larvae showed more rapid lethality compared to adult flies, despite comparable levels of Gal-1-P. The authors concluded that this observation might result from stage-specific ways in which the fruit flies responded to high Gal-1-P levels. Moreover, movement defects were demonstrated in adult flies, which did not exacerbate by early galactose exposure, despite a 20-fold increase in Gal-1-P. Galactose exposure was sufficient to cause elevated levels of Gal-1-P, without impacting the larval survival or adult outcome.

More recently, a zebrafish (Danio rerio) model of CG was created resembling the human phenotype of neurological sequel and subfertility, despite the lack of exposure to exogenous galactose. High levels of Gal-1-P were found in 9-day old larvae in the absence of exogenous galactose. These results further support the evidence that neurological and fertility impairments occur in zebrafish without exposure to exogenous galactose.

### 3.3.3 GALE deficiency

Studies in GALE-deficient S. cerevisiae demonstrated accumulated Gal-1-P levels and growth arrest following galactose exposure. The role of Gal-1-P as mediator of galactose sensitivity was studied in GALE- as well as GALT-deficient strains. GALE-deficient cells were 10-fold more sensitive to galactose compared to GALT-deficient cells and showed to remain constant Gal-1-P levels after galactose exposure, while in GALT-deficient cells the Gal-1-P levels gradually decreased over time. Loss of GALT was able to rescue the galactose-induced growth restriction of GALE-deficient cells, without impacting the Gal-1-P levels. GALE-deficient yeast also showed increased levels of UDP-Gal and slightly decreased levels of UDP-Glc, in parallel to Gal-1-P accumulation. Authors concluded that Gal-1-P alone could not explain the differential growth rates observed in galactose-sensitive yeast.

Taken together, the results from different cellular and animal models suggest that Gal-1-P is not the sole pathophysiological agent responsible for the phenotype observed in GALT and GALE deficiency. Most probably a combination of events contributes to the observed human phenotype. We hypothesize that Gal-1-P is necessary but not sufficient in all tissues for expression of the human phenotype.

### 3.4 UDP-hexoses abnormalities and glycosylation impairments

#### 3.4.1 GALT deficiency

Several reactions are involved in maintaining UDP-hexoses levels. There is evidence that in the absence of GALT, when UDP-Gal levels are low, the GALE reaction shifts toward the formation of UDP-Gal. Furthermore, UGP is responsible for the GALT-independent formation of UDP-Glc from Glc-1-P and UTP. However, UGP can also catalyse the conversion of Gal-1-P and UTP to form UDP-Gal. Additionally, studies in mice showed that UDP-Gal can possibly be formed by UGP in the absence of GALT activity.

UDP-hexose levels were studied in S. cerevisiae expressing patient GALT mutations, erythrocytes and cultured fibroblasts from patients with GALT deficiency. Yeast strains with the lowest levels of GALT activity accumulated intracellular UDP-Gal and UDP-Glc levels.
| Cell line                  | Reference          | GALT activity     | Measurement method | UDP-hexoses levels/ratios<sup>b</sup>                                                                 |
|---------------------------|--------------------|-------------------|--------------------|-------------------------------------------------------------------------------------------------------|
| Erythrocytes              | Ng et al<sup>6</sup> | Undetectable      | Enzyme-based method| Decreased UDP-Gal levels<br>• 35% of normal mean<br>Normal UDP-Gal levels<br>                                |
|                           | Berry et al<sup>51a</sup> | Absent/markedly reduced | HPLC                | Decreased UDP-Gal levels<br>• Children: 62% of normal mean, 6/19 < 95% CI for normal<br>Adults: 46% of normal mean, 4/5 < 95% CI for normal<br>Higher UDP-Glc to UDP-Gal ratio<br>• Children: 12/19 > 95% CI for normal<br>Adults: 4/5 > 95% CI for normal<br>                                                                 |
|                           | Keevill et al<sup>52</sup> | Negligible        | HPLC                | Decreased UDP-Gal levels<br>• 87% of normal mean<br>Higher UDP-Glc to UDP-Gal ratio<br>                     |
|                           | Gibson et al<sup>53</sup> | Absent/markedly reduced | HPLC                | Decreased UDP-Gal levels<br>• Children: 70% of normal mean, 10% < 2 SD of normal mean<br>Adults: 81% of normal mean<br>Higher UDP-Glc to UDP-Gal ratio<br>• 65% of children >2SD of normal mean<br>20% of adults >2 SD of normal mean<br>                                                                 |
|                           | Xu et al<sup>54</sup> | Undetectable      | HPLC                | Decreased UDP-Gal levels<br>• 62% of normal mean<br>Higher UDP-Glc to UDP-Gal ratio<br>                     |
|                           |                   | 0.05-2.5 μmol/h/g Hb (normal >20) |                   | Normal UDP-Gal levels<br>115% of normal mean<br>Normal UDP-Glc/UDP-Gal ratio<br>                          |
| Lymphoblasts              | Gibson et al<sup>55</sup> | Unknown           | HPLC                | Normal UDP-Gal levels<br>108% of normal mean<br>                                                           |
| Fibroblasts               | Ng et al<sup>6</sup> | Undetectable      | Enzyme-based method | Decreased UDP-Gal levels<br>36% of normal mean<br>Normal UDP-Gal levels<br>                               |
|                           | Gibson et al<sup>55</sup> | Unknown           | HPLC                | Normal UDP-Gal levels<br>95% of normal mean<br>Normal UDP-Glc/UDP-Gal ratio<br>                        |
|                           | Xu et al<sup>54</sup> | Undetectable      | HPLC                | Decreased UDP-Gal levels<br>55% of normal mean<br>Higher UDP-Glc to UDP-Gal ratios<br>                   |
| SV40-transformed fibroblasts | Lai et al<sup>5</sup> | Undetectable      | Enzyme-based method | Decreased UDP-Gal<br>30% of normal mean<br>Decreased UDP-Glc<br>67% of normal mean<br>Higher UDP-Glc/UDP-Gal ratio<br> |

Abbreviations: CI, confidence intervals; HPLC, high-performance liquid chromatography; UDP-Gal, uridine diphosphate-galactose; UDP-Glc, uridine diphosphate-glucose.

<sup>a</sup>Absolute levels four times lower compared to values reported by Ng et al.<sup>6</sup>

<sup>b</sup>In GALT-deficient cell lines compared to cell lines from normal controls.
after galactose exposure, possibly due to the UGP-mediated conversion of accumulated Gal-1-P to UDP-Gal. Given that the increase in UDP-Gal levels parallel those in UDP-Glc, the authors concluded that a more likely explanation might be the epimerization of the accumulated UDP-Glc to UDP-Gal by GALE, in absence of GALT activity.50

UDP-hexoses levels measured in patient erythrocytes, lymphocytes, and normal or transformed fibroblasts show sometimes contradictory results (Table 1).51-55 Drawing any conclusion from the observed findings is therefore very difficult. Various techniques (high-performance liquid chromatography [HPLC], indirect enzyme-based techniques) have been applied to measure UDP-hexoses levels, some of which are now recognised as susceptible to interference. In addition, different cell lines might behave differently and give distinct results. Furthermore, pre-analytical conditions, such as culturing and incubation techniques can also affect the results.

**Table 2** Summary of the proposed pathophysiological mechanisms in systematically reviewed studies

| Type of galactosemia | Pathophysiological mechanism | Animal/cellular model | References |
|----------------------|------------------------------|-----------------------|------------|
| GALK deficiency      | Galactitol accumulation      | Mouse                 | Ai et al26 |
|                      | Accumulation of metabolites  |                       |            |
|                      | (galactose, Gal-1-P,         |                       |            |
|                      | galactitol, galactonate)     |                       |            |
|                      | Saccharomyces cerevisiae     |                       | Lai et al5 |
|                      | Mouse                        |                       |            |
|                      | Drosophila melanogaster      |                       |            |
|                      | Danio rerio                  |                       |            |
|                      | Saccharomyces cerevisiae     |                       | Chhay et al50 |
| Abnormalities in UDP-Gal and UDP-Glc levelsa |                       |                       |            |
|                      | Patient cell lines           |                       | Dobbie et al7; Ornstein et al8; Coss et al8;                           |
|                      | (fibroblasts)                |                       | Staubach et al26 |                                      |
|                      | Patient cell lines           |                       | Lai et al5 |
|                      | (SV40-transformed fibroblasts) |                   |            |
|                      | Patient cell lines           |                       | Petry et al57 |
|                      | (lymphoblasts)               |                       | Jumbo-Lucioni et al58,59 |
| Down-regulation of PI3K/Akt pathway | Mouse                        |                       | Balakrishnan et al60,61; Chen et al62 |
| ER stress/Unfolded protein response | Saccharomyces cerevisiae     |                       | Slepak et al63; De-Souza et al64 |
|                      | Patient cell lines           |                       | Slepak et al63; Staubach et al26 |
|                      | (fibroblasts)                |                       | Balakrishnan et al51; Chen et al62 |
| Oxidative stress     | Drosophila melanogaster      |                       | Jumbo-Lucioni et al66,67 |
| Structural/functional defects | Mouse                        |                       | Tang et al50 |
|                      | Saccharomyces cerevisiae     |                       | McCorvie et al68 |
|                      | Escherichia coli             |                       | Coelho et al22 |
| GALE deficiency      | Gal-1-P accumulation         | Saccharomyces cerevisiae | Wohlers and Fridovich-Keil45; Ross et al33; Mumma et al46 |
| Abnormalities in UDP-hexoses levels | IdlD cells                   |                       | Schulz et al69 |
|                      | Saccharomyces cerevisiae     |                       | Ross et al33 |
|                      | Caenorhabditis elegans       |                       | Brokate-Llanos et al29 |
|                      | Drosophila melanogaster      |                       | Daenzer et al72; Jumbo-Lucioni et al59 |
| ER stress            | Caenorhabditis elegans       |                       | Brokate-Llanos et al29 |
| Structural/functional defects | Saccharomyces cerevisiae     |                       | McCorvie et al71 |
|                      | Escherichia coli             |                       | Pey et al72 |

Abbreviations: ER, endoplasmic reticulum; GALE, uridine diphosphate-galactose 4’-epimerase; GALK, galactokinase; UDP-Gal, uridine diphosphate-galactose, UDP-Glc, uridine diphosphate-glucose.

aA separate table (Table 3) is provided for UDP-hexoses studies in different patient cell lines.
Despite the fact that some groups report significantly lower UPD-Gal and UDP-Glc levels in erythrocytes and fibroblasts of CG patients (Table 1), the clinical relevance of the reduced levels remains questionable. Furthermore, UDP-hexoses levels in the relevant target tissues of damage (brain and gonads) have not been studied. More studies using accurate cutting-edge techniques to measure UDP-hexoses preferably in these relevant tissues of damage would greatly improve our knowledge regarding the UDP-hexoses levels in CG and its potential clinical relevance. The discrepancies in the data available hitherto do not unambiguously answer the question whether UDP-hexoses alterations are relevant for the pathophysiology and warrant further investigation.

There is evidence of abnormal glycosylation in CG patients. Haberland et al described abnormal patterns of glycoproteins in autopsy brain tissue and suggested that this could be due to defective galactose incorporation.79 Studies in fibroblasts deriving from GALT-deficient patients showed an aberrant glycosylation of glycoproteins and glycolipids.7,8 Abnormal galactose to mannose and sialic acid plus galactose to mannose ratios were found. Sialic acid is a terminal residue and is normally attached to galactose within the oligosaccharide units of glycoproteins.7 Lai et al showed a reduced abundance of the high molecular weight Sambucus nigra agglutinin (SNA) positive glycoproteins in GALT-deficient SV40-transformed fibroblasts exposed to galactose, indicating an abnormal glycosylation.5

Coss et al demonstrated dys-regulation of the glycosylation and inflammatory pathways at the gene and protein level in GALT-deficient fibroblasts.4 Imbalance in glycolipids in GALT-deficient lymphoblasts was found by Petry et al, hypothesized to be caused by deficient levels of UDP-Gal and UDP-GalNAc and thereby reduction of galactosylation products.57 In addition, Staubach et al showed a decrease in plasma membrane expression of N-glycoproteins upon galactose exposure in GALT-deficient fibroblasts, important in the regulation of cell signalling. Authors suggested that at least for some of the glycosylated membrane proteins a dys-glycosylation might be the cause of decreased membrane expression.56

Jumbo-Lucioni et al studied the synaptomatrix glycosylation in the GALT-deficient D. melanogaster and showed that glycans play a primarily inhibitory role in modulating synapse morphogenesis. They suggested that neuromuscular junction development defects and associated impairments in coordinated movement in the GALT-deficient fruit flies might result from disrupted UDP-hexose balance and the consequently impaired glycosylation.58 Double mutant GALT-, UGP-deficient fruit flies showed exacerbated coordinated movement deficits. UGP played a particularly important role enabling coordinated movement and was found to be a strong negative modulator of neurobehavioral outcomes in the fruit flies.59
3.4.2  |  GALE deficiency

The GALE deficiency was studied in the cellular models, *S. cerevisiae*, the *Caenorhabditis elegans* and the *D. melanogaster* models.

GALE-deficient *S. cerevisiae* accumulated UDP-Gal and showed decreased levels of UDP-Glc upon galactose exposure.

Schulz et al showed that GALE-deficient ldlD cells (GALE-impaired mammalian cells utilising a strain of Chinese hamster ovary (CHO)) accumulated both Gal-1-P and UDP-Gal when exposed to galactose, whereas depleted levels of UTP, UDP-Glc, UDP-GalNAc, and UDP-GlcNAc were found. An explanation given by the authors is that absent of GALE activity prevented the production of UDP-Gal into UDP-Glc. The accumulation of UDP-Gal resulted in a UDP-sink, leading to UTP/uridine depletion. Because UTP is necessary for the de novo synthesis of UDP-Glc by UGP and UDP-GlcNAc by UDP-N-acetylglucosamine pyrophosphorylase, the cells were not able to synthesise these products. Therefore, UTP depletion might contribute to the galactose sensitivity in these cells. Accumulation of Gal-1-P and UDP-Gal correlated with galactose-induced compromised growth of GALE impaired cells, whereas deficiency of UDP-GalNAc did not. Cells expressing *E. coli* GALE exhibiting activity exclusively toward UDP-Gal/UDP-Glc (lacking activity toward UDP-GalNAc/UDP-GlcNAc), grew normally in the presence of galactose, despite undetectable levels of UDP-GalNAc.

Studies in the GALE-deficient *C. elegans* model showed increased levels of UDP-Gal and reduction of UDP-GalNAc. The reduction of GALE activity affected multiple aspects of development, most likely resulting from a general defect in the glycosylation process. Strong developmental defects were seen in gonad migration and vulva formation, possibly resulting from a decrease in the developmental defects were studied and showed that both activities were necessary for survival. Loss of activity toward UDP-Gal was shown to result in Gal-1-P and UDP-Gal accumulation, a reduced lifespan when exposed to galactose and impaired fecundity in both male and female larvae, whereas loss of activity toward UDP-GalNAc did not lead to metabolic abnormalities but resulted in compromised survival and impaired fecundity in female larvae. Authors suggested that nucleotide sugars imbalance rather than Gal-1-P was responsible for the compromised survival and fecundity in this model.

Jumbo-Lucioni et al showed that loss of neuronal GALE activity worsened the coordinated movement defects in GALT-deficiency. GALE-deficient fruit flies showed compromised coordinated movement as well as neuromuscular junction synaptogenesis defects. Furthermore, the GALE deficient fruit flies showed also loss of UDP-GalNAc in the synaptomatrix.

The results provided by these cellular and animal studies reveal UDP-hexose impairments in GALE deficiency. The degree of severity of the clinical phenotype depends on the impairment of GALE activity (toward UDP-Gal/UDP-Glc and/or UDP-GalNAc/UDP-GlcNAc).

4  |  SIGNALLING PATHWAYS ALTERATIONS, UPR ACTIVATION, ER STRESS, AND OXIDATIVE STRESS

4.1  |  GALT deficiency

Signalling pathway alterations have been investigated in GALT-deficient mouse fibroblasts and isolated ovaries and cerebella from GALT-deficient mice. These studies demonstrated that the phosphatidylinositol-4,5-bisphosphate 3-kinase (PI3K)/protein kinase B (Akt) signalling pathway was down regulated. This pathway is essential for the growth, proliferation, and survival of many tissues, including neuronal and reproductive tissues. Studies in GALT-deficient mouse fibroblasts showed that the down regulation of the PI3K/Akt pathway occurred at both mRNA and protein levels. The authors proposed ER stress and aberrant glycosylation as candidate factors to connect GALT deficiency and the signalling pathway alterations.

The increase in binding protein (BiP) abundance and the up regulation of *Hspb1* that were found, indicated ER stress. This ER stress resulted in reduced mRNA level of *pan-Akt* and its abundance, which could lead to decreased Akt availability for phosphorylation. Moreover, a decreased *pan-Akt* level and diminished *pAkt*(thr308) and *pAkt* (*Ser473*) phosphorylation were found.

Studies by Chen et al and Balakrishnan et al showed a down-regulated PI3K/Akt signalling pathway in the cerebellum isolated from GALT-deficient mice. They found a significantly reduced steady-state protein abundance of *pAkt* (*Ser473*), *pAkt* (Thr308), *pan-Akt*, and pGSK-3β. Furthermore, an increased abundance of BiP was found in this tissue. Decreased thicknesses of the molecular and granular layer in the cerebellum of mutant mice suggested cerebellar hypoplasia. These mice manifested ataxia-related motor impairments with varying degrees of severity. A progressive nature of the impairments in older mice was observed. Additionally, studies by Balakrishnan et al showed down-
regulated of the PI3/Akt pathway in isolated GALT-deficient ovaries. pAkt (Ser473), pGsk3β, and Hsp90 levels in the isolated GALT-deficient ovaries were significantly reduced and higher levels of BiP were found. Moreover, less primordial follicles and Purkinje cells were seen in GALT-deficient mice.61,82 The PI3/Akt signalling is important in supporting the survival of immature ovarian follicles.82

ER stress and the unfolded protein response (UPR) activation were studied in GALT deficient *S. cerevisiae* and fibroblasts.

In mammalian cells, the UPR signal transduction pathway is designed to cope with ER stress, caused by accumulation of misfolded proteins in the ER.83-85 If the UPR activation is prolonged, it can trigger pro-apoptotic pathways.86,87 Slepak et al showed that accumulated levels of Gal-1-P exerted down regulation of ribosomal protein (RP) genes (eg, RPA12 and RRN11) and genes involved in the various aspects of RNA metabolism (eg, UTP6, SNR54 snRNA) in GALT-deficient yeast exposed to galactose.63 De-Souza et al showed that Gal-1-P synthesis was essential to induce ER stress by galactose and consequently UPR activation in a yeast model of CG. A defect in inositol metabolism and subsequently calcium homeostasis was proposed to contribute to the observed ER stress.64

Studies in human GALT-deficient fibroblasts exposed to galactose showed that the accumulated Gal-1-P competitively inhibited human inositol monophosphatase, leading to inositol depletion. Furthermore, they showed that as a result, Ca²⁺ was released from the ER.65 Perturbation of calcium homeostasis in the ER is known to be an initiator of ER stress.85 Slepak et al found higher expression of ER stress related genes in GALT-deficient fibroblasts (CHOP, calreticulin, BiP, GRP94, and XBP-1) after galactose exposure.65

Staubach et al found increased levels of BiP and calreticulin in GALT-deficient fibroblasts after galactose exposure.56 They are stress markers that indicate activation of the UPR and are involved in protein folding (GRP78) and protein quality control of N-glycoproteins by binding dysglycosylated molecules (calreticulin).88 Sustained activation of the UPR has been implicated in the progression of several neurodegenerative diseases.89

Moreover, studies in *D. melanogaster* revealed that oxidative stress and the response to stress contributed to the mechanism of acute galactose toxicity in GALT deficiency. Galactose exposure led to oxidative stress in both GALT-deficient and control fruit fly, but they responded differently. An induction of genes involved in glutathione metabolism and known to function in the response to oxidative stress (GSTD6 and GSTE7)90 were found in both GALT-deficient and control larvae, however, GALT-deficient larvae showed diminished response. The survival rates of galactose-exposed mutant animals were affected by the presence of oxidants and antioxidants whereas the survival rates of control animals were unaffected.66 In 2014, Jumbo-Lucioni et al demonstrated that GALT-deficient larvae exposed to galactose showed increased protein oxidative damage, with almost twice the level of protein-bound cysteine and glutathione compared to controls.67

Above presented results suggest that ER stress plays an important role in the pathophysiology of GALT deficiency. Subsequently, ER stress can induce UPR and signalling pathway alterations. Furthermore, oxidative stress also seems to be of influence in the observed phenotype.

### 4.2 | GALE deficiency

The expression of *C. elegans* genes involved in the activation of the UPR pathway (*hsp-4* and *xbp-1*) were altered in this GALE-deficient animal model, thus inducing ER stress. Authors suggested that the observed ER stress might be caused by the incorrect glycosylation of proteins involved in ER functions or by alterations of the UDP-Glc/UDP-Gal ratio.29

### 4.3 | Structural and functional defects

### 4.4 | GALT deficiency

Several studies in *S. cerevisiae* and *E. coli* have shown that mutations in the human GALT gene resulted in a less stable variant of the protein which failed to fold correctly, was more susceptible to proteolysis or formed aggregates.22,68 In particular, the p.Q188R variant (most common *GALT* mutation variant in the Caucasian population) affect GALT aggregation propensity.22 The yeast model (described by21) has been used to study the critical role of different mutations in determining the expression and functionality of the human GALT enzyme.50,91-94

### 4.5 | GALE deficiency

McCorvie et al showed that altered cofactor binding can affect stability and activity of human GALE in an *E. coli* and *S. cerevisiae* expression systems. A certain level of flexibility was shown to be required for proper function. An increase or decrease in flexibility of GALE, as seen with p.K161 N and p.D175 N, respectively, resulted in loss of function.71 Studies by Pey et al concluded that
destabilisation and altered conformational dynamics of GALE dimer at physiological conditions (temperature and pH) were linked to loss of enzyme function in GALE deficient *E. coli*. The authors found that natural ligands (NAD+ or UDP-Glc) could stabilise the protein.72

In conclusion, in both GALT- and GALE-deficiency, numerous disease-causing mutations are described be give rise to variants with higher propensity for aggregation and altered stability.

### 4.6 | Treatment approaches

Since the description of the first infant with hypergalactosemia who responded well to a lactose-restricted diet,95 galactose restriction is the cornerstone of treatment for the galactosemias. The understanding of the molecular basis underlying this group of diseases hitherto yields several other treatment approaches possibilities.

#### 4.6.1 | GALK inhibition

Inhibition of GALK, as a potential therapeutic strategy, has been studied. In GALT-deficient yeast secondary deletion of *GALK1* stabilised the levels of UDP-Gal and UDP-Glc and relieved the galactose-induced lethality in both GALT- and GALE-deficient cells.33 Furthermore, it alleviated the galactose-mediated growth arrest of GALT- and GALE-deficient yeast.46 *GALK1* deletion completely inhibited the galactose-dependent UPR activation and galactose toxicity in yeast,64 but did not increase tolerance to other ER stressors such as tunicamycin or dithiothreitol. Furthermore, GALK inhibitors had a positive influence on ER stress and on the phosphate levels in GALT-deficient yeast.73 The effect of GALK inhibitors in GALT-deficient fibroblasts showed a dose-dependent effect on Gal-1-P accumulation, where levels were lowered to control levels. Tang et al provided a proof-of-concept that GALK inhibition can lower Gal-1-P in galactose-challenged GALT-deficient fibroblasts.74

GALK inhibition was studied in the *D. melanogaster*. Jumbo-Lucioni et al showed that *GALK1* co-removal corrected the movement defects observed in GALT-deficient fruit flies and restored the synaptomatrix glycosylation state, however, it did not fully reverse NMJ glycan changes.58 Deletion of *GALK1* in the GALT-deficient fruit flies was studied further and revealed to be beneficial in reducing the levels of Gal-1-P, but not in rescuing survival of larvae to adulthood, motor defects or fecundity. On the contrary, deletion of *GALK1* had a significant negative impact on climbing ability and fecundity in GALT-deficient fruit flies, in the absence of galactose.34

Taken together, GALK inhibitors are effective in rescuing the biochemical phenotype (eg, reducing Gal-1-P levels), however, the effect on the clinical phenotype remains not fully clear. Systematically summarising the current evidence in galactosemia’s pathophysiology revealed that Gal-1-P is not the sole pathogenic agent responsible for the observed disease complications.

#### 4.6.2 | Antioxidants

Increased oxidative stress has been reported in the GALT-deficient *D. melanogaster*. The antioxidants vitamin C and alpha-mangostin increased the survival rates of mutant larvae to adulthood, whereas Gal-1-P remained elevated, suggesting that these antioxidants act downstream or independently of Gal-1-P.66 In addition, free radical scavengers that mimic the action of Mn2+-based superoxide dismutase (SOD) have been shown to be effective in rescuing larval survival and reverse protein oxidative damage in GALT-deficient flies in the presence of galactose.67 Reactive oxygen species (ROS), scavenged by the SOD mimics, contributed to the acute galactose toxicity and galactose-independent long-term complications in GALT-deficient adult fruit flies. These SOD mimics are described to accumulate in critical intracellular compartments, including mitochondria, and promote binding to electron-rich anionic reactive species enabling them to scavenge superoxide, peroxynitrite, and other free radicals.96 These compounds showed little toxicity, and thus have considerable potential as lead compounds for therapeutic strategies.67

Antioxidant therapy could be one of the possible co-adjuvant therapies that needs further evaluation.

#### 4.6.3 | Pharmacological chaperones

In 2016, the crystal structure of human GALT was resolved97 that provided insights into the functional impact of mutations. The observation that many of the disease-causing mutations of human *GALT* and *GALE* showed structural instability has led to the suggestion that pharmacological chaperones that stabilise and increase activity of the affected protein could be beneficial.68,72 The effect of arginine, a well-established suppressant of protein aggregation,98 has been studied in an *E. coli* model. Arginine has shown a mutation-specific effect, with rescue of human GALT p.Q188R, p.K285 N, and p.G175D variants, suggesting that this amino acid could be of therapeutic benefit in patients carrying these mutations.73 However, studies in GALT-deficient fibroblasts from patients carrying the p.Q188R exposed to arginine did not show changes in GALT activity. Thermal shift analysis of recombinant p.Q188R GALT protein in
the presence of arginine did not exhibit a positive effect. The authors also evaluated this approach in four p. Q188R homozygous patients but it did not prove effective.76

4.6.4 | Antisense therapy

Coelho et al employed two locked nucleic acid oligonucleotides to successfully restore the aberrant splicing profile caused by the intronic mutation c.820 + 13A > G (IVS8 + 13A > G), thus, establishing a proof of concept for the application of antisense therapy for mis-splicing mutations in CG.77

4.6.5 | ER stress reducers

Balakrishnan et al showed that diminished PI3K/Akt signalling, induced by ER stress, played an important role in subfertility and cerebellar ataxia in the GALT-deficient mouse. Follicle loss due to insufficient Akt activity could be responsible for depletion of most of the primordial follicle pool very early in postnatal life. The effect of salubrinal, an experimental ER stress reducer, was evaluated in the mice. Treatment with salubrinal increased the numbers of primordial follicles. Furthermore, it significantly slowed down the loss of Purkinje cells in GALT-deficient mice.60,61

The chemical chaperone 4-phenylbutyric acid was not effective in reducing ER stress in a S. cerevisiae model for CG.64 De-Souza et al suggested the idea that molecules that interfere with ER stress might be good drug candidates to treat CG.

4.6.6 | UGP up-regulation

Lai and Elsas showed that UGP1 up-regulation in isolated GALT revertant from S. cerevisiae cells succeeded to help overcome the galactose toxicity in the GALT revertant, reduced the level of Gal-1-P and could replenish UDP-Gal. The over-expression of the human homologue hUGP2 gene alone could rescue the growth defect of the GALT-deficient S. cerevisiae exposed to galactose.78 Studies in GALT-deficient SV40- transformed fibroblasts showed increased UDP-hexoses levels and no accumulation of Gal-1-P in cells expressing human UGP2.5

Based on the studies hitherto described, human UGP2 up-regulation could be a candidate as therapeutic approach and should also be evaluated in animal models.

4.6.7 | Uridine supplementation

The effect of uridine was evaluated in erythrocytes from GALT-deficient patients and demonstrated that UDP-Glc and UDP-Gal levels increased to control levels upon supplementation, without impacting the Gal-1-P level. When exposed to both uridine and galactose, UDP-Gal increased even more compared to uridine alone. The fact that uridine increased UDP-Gal levels in the absence of GALT activity suggested another mechanism, possibly through the UGP reaction.5

In GALE-deficient ldlD cells, uridine supplementation relieved the galactose-specific growth inhibition and corrected UDP-Glc and to a lesser extent UDP-GalNAc and UTP, without resucing the Gal-1-P and UDP-Gal levels. A large pool of UDP-Gal leading to uridine depletion and a potential spectrum of negative downstream effects in the GALE-impaired cells exposed to galactose was suggested as a key mediator in GALE-impaired mammalian cells.69 In 1997, Manis et al studied the effects of oral uridine supplementation (150 mg/kg/day for a mean of 3.1 years) in 35 patients (aged 1 week-16.2 years) with GALT deficiency. Uridine supplementation did not prove to be effective in improving cognitive functioning in these patients compared with patients that were only on a galactose restricted diet.99

Taken together, results obtained from cellular studies and the clinical trial in patients with GALT deficiency show that uridine supplementation is not promising in rescuing the biochemical or clinical phenotype observed in patients.

This study has several limitations. First, we excluded original articles that were not written in English. Also, we have excluded the hypergalactosemia models from this study and focused only on hereditary galactosemia, thereby possibly missing insights on the pathophysiology that underlies galactosemia. Furthermore, results obtained from the cellular and animal models cannot directly be extrapolated to the human situation. The strengths of this study include the comprehensive search and a large number of included studies. Moreover, the inclusion and data extraction have been performed by independent researchers, thereby reducing the risk of bias. To the best our knowledge, this is the first study that provides a comprehensive overview of the scattered information regarding the pathophysiology of hereditary galactosemia based on studies in cellular and animal models performed in the past decades.
This information can be of great benefit to researchers in the field and provides insights on potential treatment targets and approaches.

5 | CONCLUSIONS

Cellular and animal models have considerably contributed to the pathophysiological insights underlying hereditary galactosemia. Results obtained from this systematic review of the hitherto performed cellular and animal studies reveal that Gal-1-P is not sole pathophysiological agent responsible for the phenotype observed in GALT and GALE deficiency. We hypothesize that Gal-1-P is necessary but not sufficient in all tissues for expression of the human phenotype. Other currently described contributing factors that underlie the pathophysiology of acute and long-term outcomes in GALT and GALE deficiency include galactose metabolites accumulation, UDP-hexoses alterations and subsequent impaired glycosylation, ER stress, signalling pathway alterations and oxidative stress.

ER stress plays an important role in the pathophysiology of GALT deficiency. Subsequently, ER stress can induce UPR and signalling pathway alterations. Furthermore, oxidative stress is another contributing factor in the observed phenotype. Structural and functional studies show that both GALT and GALE disease-causing mutations give rise to variants with a higher propensity for aggregation and altered stability. The crystal structure of human GALT that was resolved in 2016, allows us to study new therapies enhancing GALT activity/stability, including pharmacological chaperones.

Several therapeutic approaches addressing the contributing factors in the pathophysiology of hereditary galactosemia have been subsequently evaluated in cellular and animal models. GALK inhibitors are found to be effective in rescuing the biochemical phenotype (eg, reducing Gal-1-P levels), however, the effects on the observed clinical phenotype remain unclear. Other treatment approaches that have shown to rescue the biochemical and/or clinical outcome include antioxidants, pharmacological chaperones, ER stress reducers, UGP up-regulation, and uridine supplementation. Based on the hitherto performed studies, possible co-adjuvant therapies that need further evaluation include antioxidant therapy and UGP up-regulation. Results from cellular studies and studies in patients show that uridine supplementation is not promising in rescuing the biochemical phenotype. Studies investigating the utilisation of the optimum therapeutic window are crucial for the success of any treatment modality.

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CONFLICT OF INTEREST

The authors declare no potential conflict of interest.

AUTHOR CONTRIBUTIONS

M.H. is contributed in conception and design, acquisition of data, analysis and interpretation of data, drafting the article and revising it critically for important intellectual content. A.I.C. is contributed in conception and design, acquisition of data, analysis and interpretation of data, revising the article critically for important intellectual content. J.B. is contributed in revising the article critically for important intellectual content. J.M.V. is contributed in revising the article critically for important intellectual content. L.K.M.S. is contributed in revising the article critically for important intellectual content. G.T.B. is contributed in conception and design, revising the article critically for important intellectual content. M.E.R.-G. is contributed in revising the article critically for important intellectual content. E.V.-M. is contributed in conception and design, revising the article critically for important intellectual content. L.J.I.Z. is contributed in revising the article critically for important intellectual content. A.I.C. is contributed in conception and design, acquisition of data, analysis and interpretation of data, drafting the article and revising it critically for important intellectual content. M.H. is contributed in conception and design, acquisition of data, analysis and interpretation of data, drafting the article and revising it critically for important intellectual content. A.I.C. is contributed in conception and design, acquisition of data, analysis and interpretation of data, drafting the article and revising it critically for important intellectual content. M.H. is contributed in conception and design, acquisition of data, analysis and interpretation of data, drafting the article and revising it critically for important intellectual content. A.I.C. is contributed in conception and design, acquisition of data, analysis and interpretation of data, drafting the article and revising it critically for important intellectual content. M.H. is contributed in conception and design, acquisition of data, analysis and interpretation of data, drafting the article and revising it critically for important intellectual content. A.I.C. is contributed in conception and design, acquisition of data, analysis and interpretation of data, drafting the article and revising it critically for important intellectual content.

ORCID

Minela Haskovic https://orcid.org/0000-0003-2685-6811

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

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