Factors Affecting Compliance to a Gluten-Free Diet in Pediatric Populations with Celiac Disease

Vítor Macedo-Campos¹,²,³,⁴,§ Rui Macedo-Campos¹,²,³,⁴,§ Filipa Pinto-Ribeiro¹,³,⁴ Henedina Antunes¹,²,³,⁴

¹ School of Medicine, University of Minho, Braga, Portugal
² Gastroenterology, Hepatology and Nutrition Unit, Pediatric Department and Clinical Academic Center, Hospital de Braga, Braga, Portugal
³ Life and Health Sciences Research Institute (ICVS), School of Medicine, University of Minho, Braga, Portugal
⁴ ICVS/3B’s—PT Government Associate Laboratory, Braga-Guimarães, Portugal

Address for correspondence Vítor Manuel Macedo-Campos, Rua das Portelas, N342, Góios Barcelos 4755-248, Portugal (e-mail: vitormaccampos@hotmail.com).

Abstract
Celiac disease (CD) is a multisystemic autoimmune disorder triggered by gluten, and the only known remedy available for this malady is a gluten-free diet (GFD). Therefore, we performed a systematic review to correlate the influence of different factors in compliance to a GFD. We searched PubMed database, from inception to April 2019. As inclusion criteria we considered population under 18 years, confirmed diagnosis of CD without related comorbidities and the study objective being the factors affecting compliance to a GFD. The variables compared were age, parent’s education level, parental knowledge about CD, family type, celiac association membership, quality of life, and perception of difficulties in maintaining a GFD. We identified 1,414 articles, 35 articles were eligible for full text assessment and 12 were included in the study since they studied similar variables. Our work has found some limitations namely a variety of methods to assess GFD compliance, a limiting definition of compliance, a parental bias in data, an absence of standardization in age categories, and a majority of studies being observational in their nature. Age as well as parental knowledge of CD and family type are key factors in pediatric GFD compliance. Nevertheless environmental, social, and family factors were also related with compliance. Further studies are needed to fully disclose the causality relation between these factors and compliance.

Keywords
► celiac disease
► gluten-free diet
► adherence
► pediatric
► systematic review

Introduction
The prevalence of celiac disease (CD) in Europe is approximately 1%.¹ The pathophysiology for CD involves a multisystemic autoimmune-mediated disorder to gluten, a protein most commonly found in wheat, rye, and barley resulting in injury to the small bowel mucosa.² Genetic susceptibility with HLA DQ-2 and/or DQ-8 positivity is strongly associated with the disease.²

Diagnosing CD is challenging due to its nonspecific and heterogeneous clinical presentation. The symptoms can vary in intensity but commonly it presents with abdominal symptoms such as malabsorption, discomfort, loose stools, and flatulence³ and a variety of nonintestinal symptoms that include short stature, infertility, delayed puberty, anemia, liver abnormalities, joint and muscular disorders, neurological complications, psychiatric disorders and cutaneous and mucosal manifestations.⁴ Importantly, CD can also affect asymptomatic patients.³

§ These authors contributed equally to this work.
Regarding the treatment of CD, the only knowledgeable efficient treatment is a gluten-free diet (GFD). Without the exposure to gluten, the symptoms as well as the damage inflicted, regress and the patient becomes asymptomatic. Nevertheless, it represents a major lifelong change in lifestyle. Compliance with GFD can be challenging, onerous, expensive, and imposes difficulties to the patient. Hence, the need for a systematic review to identify factors that interfere with GFD compliance and to recognize predictors of noncompliance as well as modifiable factors that positively influence compliance is fully justified.

Methods

Search Strategy
We searched the PubMed database for literature regarding the compliance to GFD in pediatric CD from inception to April 16, 2019. The terms used to perform the search were as follows: celiac or celiac or gluten sensitive enteropathy AND diet$ or nutrition$ or GFD or gluten-free or gluten free AND advice or adherence or compliance or concordance or prescription or intervention or management AND child$ or pediatric$. No filter was applied.

Eligibility Criteria for Studies and Participants
The inclusion criteria were a study population that included parents of or children under 18 years old throughout the entire study course; confirmed diagnostic of CD without related comorbidities; and focus on the factors influencing compliance to a GFD.

The exclusion criteria were articles not written in English and gray literature.

Study Selection and Data Extraction
Study selection was performed independently by two authors (V.M.-C. and R.M.-C.). Discrepancies were resolved through discussion among them or by consulting a third author (H.A.). A PRISMA flowchart (Fig. 1) was used to perform this record.

Prior to data extraction, the authors through the analysis of the papers herein included develop an excel form to

![Flowchart representing the study selection process.](image)
systematically extract data (V.M.-C. and R.M.-C.). The form contemplated the following detailed parameters: title, authors, country of origin, year of publication, study design, participant number, mean participant age, the specific aim of study, methods used to assess compliance, and factors related with GFD compliance.

**Study Quality and Assessment of Risk Bias**

Quality was individually assessed by two authors (V.M.-C. and R.M.-C.). The authors resolved any disagreement through discussion of each dissonant parameter. When agreement could not be reached, a third author was consulted (H.A.).

Since the articles retrieved were cohort, cross-sectional, and case–control studies, the authors decided to assess quality using the National Heart, Lung and Blood Institute Study Quality Assessment Tools. This checklist comprises 12 questions for case–control studies and 14 questions for cross-sectional and cohort studies. The questions must be answered using “Yes,” “No,” or “Cannot be applied/Not answered/Not reported.”

After finalizing the quality assessment, the articles were divided into three groups rated as “Good,” “Medium,” or “Poor” according to their final scores. To obtain these subcategories, we excluded the questions in which all the articles scored the same. This information is summarized in Tables 1 and 2.

### Results

**Description of Study Selection**

The flowchart in Fig. 1 shows a schematic presentation of the selection process of studies included. After searching

| Question | Yes | No | Cannot Determine/Not Applicable/Not Reported | Medium | Poor |
|----------|-----|----|---------------------------------------------|--------|------|
| 1. Was the research question or objective in this paper clearly stated and appropriate? | | | [–] | | |
| 2. Was the study population clearly specified and defined? | | | [–] | | |
| 3. Was the participation rate of eligible persons at least 50%? | | | | | |
| 4. Were all the subjects selected or recruited from the same or similar populations (including the same time period)? | | | | | |
| 5. Was a sample size justified? | | | | | |
| 6. For the analyses in this paper, were the exposure(s) of interest measured prior to the outcome(s) being measured? | | | | | |
| 7. If less than 100% of eligible cases and/or controls were selected for the study, were the cases and/or controls randomly selected from those eligible? | | | | | |
| 8. Was there use of concurrent controls? | | | | | |
| 9. Were the investigators able to confirm that the exposure/risk occurred prior to the development of the condition or event that defined a participant as a case? | | | | | |
| 10. Were the measures of exposure/risk clearly defined, valid, reliable, and implemented consistently across all study participants? | | | | | |
| 11. Were the assessors of exposure/risk blinded to the case or control status of participants? | | | | | |
| 12. Were key potential confounding variables measured and adjusted statistically in the analyses? If matching was used, did the investigators account for matching during study analysis? | | | | | |
| 13. Was loss to follow-up after baseline 20% or less? | | | | | |
| 14. Were key potential confounding variables measured and adjusted statistically for their impact on the relationship between exposure(s) and outcome(s)? | | | | | |

### Table 1 Quality assessment of case–control studies

| Study | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 | 13 | 14 |
|-------|---|---|---|---|---|---|---|---|---|----|----|----|----|----|
| Chauhan et al | + | + | – | – | – | + | + | ? | – | + | + | + | ? | Good |
| Wagner et al | + | + | – | – | – | + | + | ? | – | + | + | + | – | Medium |
| Ljungman and Myrdal et al | + | – | – | + | + | ? | – | + | + | – | + | – | – | Poor |

| Study | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 | 13 | 14 |
|-------|---|---|---|---|---|---|---|---|---|----|----|----|----|----|
| Mager et al | + | + | + | + | – | – | – | – | + | + | ? | ? | + | Good |
| Anson et al | + | + | + | – | – | – | – | + | + | ? | – | – | – | – |
| Sarkhy et al | + | ? | + | + | – | – | + | + | – | + | ? | + | Medium |
| Charalampopoulos et al | + | + | + | – | – | – | – | + | + | ? | + | Medium |
| Taghdir et al | + | + | + | – | – | – | + | + | ? | – | – | – | – | Medium |
| Khurana et al | + | – | ? | + | – | – | – | – | + | + | + | ? | ? | Poor |
| MacCulloch and Rashid | + | + | ? | + | – | – | – | – | + | + | ? | – | – | Good |
| Garg and Gupta | + | + | ? | + | – | – | – | – | + | + | ? | + | Good |
| Roma et al | + | + | ? | + | – | – | – | – | – | + | + | ? | – | Poor |

([–], “No”; [+], “Yes”.)
PubMed using the keywords as described earlier, we obtained a total of 1,414 results. Nevertheless, 1,379 papers were excluded based on their title or abstract. Exclusion was based on the article type (systematic review, meta-analyses, comment, expert opinion, or letter), CD being associated with some comorbidity, compliance to GFD not being the main aim, and population over 18 years old. Of the 35 articles that remained for full text assessment, 23 articles were not eligible based on the following premises: seven studied population with patients older than 18 years old, five did not regard CD, four did not correlate with compliance, four presented no comparable data, two with no statistical evidence and one could not be accessed in time for this review. A total of 12 articles were finally analyzed.

Methodological Quality
Of the included studies, nine are cross-sectional (75%) and three are case–control (25%) studies. Two cross-sectional studies and one case–control article were rated as “Poor.” This rating was tied to the following assessments: Khurana et al\(^8\) lacked information regarding population details and Roma et al\(^9\) and Ljungman and Myrdal\(^10\) did not discriminate the factors affecting compliance, namely those that were evaluated as a dichotomous variable when more discrimination was possible.

The cross-sectional papers that scored the lowest were because of the following reasons: justifying and pretending the results of statistical analysis, exposure being measured before assessing the outcome, the time frame used, number of exposure assessment, number of participants lost to follow-up (not applicable to this type of study), and identification of potential confounding factors.

Regarding case–control studies, the items where all papers scored the lowest were the justification for the sample size and the use of concurrent controls.

Characterization of the Population
Data concerning the population at study is summarized in Table 3. A population of 1,579 children was included in our study. None of the papers used populations smaller than 40 patients? Four articles\(^9,10,12,14\) did not display the mean age and two described it using a median statistic.\(^14,15\) Therefore, we were able to estimate an average CD patient age of 10.3 years. The papers report to different geographic areas with five being from Europe,\(^5,10,12,14\) three from India,\(^8,11,13\) another three from Middle East,\(^5,16,17\) and two from Canada.\(^15,18\) Except for two of them,\(^10,17\) all papers were published within the last decade. Table 3 also elucidates on the method and criteria used for the diagnosis of CD.

Characterization of the Study
The analyses of Tables 4 and 5 summarize the results of each paper considering the following categories: demographic factor, household, child related, parent related, dietary related, disease related, personality related, and quality of life (QoL).

Adhesion Assessment
Adhesion was measured heterogeneously between studies. One study\(^8\) (9%) evaluated compliance measuring antitissue transglutaminase antibodies (t-TG) levels whereas 11 studies\(^5,9,10,12–16,18\) (91%) relied on self-reported information with only one verifying it through measurement of t-TG levels.\(^18\) Self-reported compliance can be divided in questionnaires\(^5,9,10,12–16,18\) or

Table 3 Characterization of the population at study and criteria for the diagnosis of CD

| Authors            | Country     | Year | Study design | Study population | Age mean ± SD | Criteria used for the diagnosis of CD | Biopsy to confirm CD |
|--------------------|-------------|------|--------------|------------------|---------------|---------------------------------------|----------------------|
| Mager et al        | Canada      | 2018 | Cross sectional | 372              | 10.4 ± 3.8 (CD) 10.9 ± 4.0 (controls) | Nondisclosed           | Performed            |
| Chauhan et al      | India       | 2010 | Case–control  | 70               | –             | ESPGHAN 2012                          | Performed            |
| Anson et al        | Israel      | 1989 | Cross sectional | 43               | 10.7          | Nondisclosed                          | Performed            |
| Sarkhy et al       | Saudi Arabia| 2015 | Cross sectional | 133              | 9.9           | Nondisclosed                          | Performed in 94% of population |
| Wagner et al       | Austria     | 2016 | Case–control  | 376              | –             | Nondisclosed                          | Performed            |
| Ljungman and Myrdal| Sweden      | 1993 | Case–control  | 47               | –             | ESPGHAN 1969                          | Performed            |
| Taghdir et al      | Iran        | 2016 | Cross sectional | 65               | 11.3 ± 3.8    | ESPGHAN criteria (year nondisclosed)  | Performed            |
| Charalampopoulos et al | Greece | 2013 | Cross sectional | 90    | 12.1 (median) | ESPGHAN 2012                          | Performed            |
| Roma et al         | Greece      | 2010 | Cross sectional | 73               | 10.4          | ESPGHAN 1990                          | Performed            |
| Khurana et al      | India       | 2014 | Cross sectional | 50               | 9.06         | ESPGHAN 1990                          | Performed            |
| MacCulloch and Rashid | Canada | 2014 | Cross Sectional | 126              | 12 (median)    | Nondisclosed                          | Performed            |
| Garg and Gupta     | India       | 2014 | Cohort        | 134              | –             | ESPGHAN 1990                          | Performed            |

Abbreviations: –, value was not disclosed; CD, celiac disease; ESPGHAN, European Society for Paediatric Gastroenterology Hepatology and Nutrition.
Table 4 Discrimination of variables regarding demographic, household, child, and parent-related factors

| Demographic factors | Household | Child Related |
|---------------------|-----------|--------------|
| Gender | Maternal age | Paternal age | Maternal education | Paternal education | Age | Non-Caucasian | Born in Canada | Family history | Residence | Parental occupational status | Family type | Higher income per capita | Less number of siblings (0-1) | Parents’ country of origin | Parents’ marital status | Parents’ labor force participation | Working or non-working mother |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| Mager et al | x | x | x | x | x | ♦ | ♦ | - | - | - | - | - | - | - | - | - | - | - |
| Chauhan et al | x | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - |
| Anson et al | x | - | - | - | - | ♦ | ♦ | - | - | - | - | - | - | - | - | x | x | x | - |
| Sarkhy et al | - | - | - | x | - | - | - | x | x | - | - | - | - | - | - | - | - | - | - |
| Ljungman and Myrdal | ♦ | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - |
| Taghdir et al | x | - | - | - | - | ♦ | ♦ | - | - | - | x | - | - | - | - | - | - | - | - |
| Wagner et al | x | - | - | - | - | - | - | x | - | - | - | - | - | - | - | - | - | - | - |
| Charalampopoulos et al | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - |
| Roma et al | - | - | - | x | x | - | - | - | - | - | - | - | - | - | - | - | - | - | - |
| Khurana et al | x | - | - | - | x | x | - | - | - | - | - | x | - | - | - | - | x | x | - |
| MacCulloch and Rashid | x | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - |
| Garg and Gupta | x | - | - | - | - | ♦ | ♦ | - | - | - | - | - | - | - | - | - | - | - | - |

| Demographic factors | Household | Child Related |
|---------------------|-----------|--------------|
| Mother outside home | Siblings without CD | Province of living | Family income per capita | Occupational status of the father (Professional) | Social support or the lack of it | Habitation | Number of individuals in the household consuming a GFD | Number of children in the household | Household income were noted | Weight | Height | Site of investigation | Birth order | Number of siblings | Child knowledge of CD | Current BMI |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| Mager et al | - | - | - | - | - | - | x | x | x | x | x | x | x | x | x | x | - | - | - |
| Chauhan et al | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - |
| Anson et al | - | - | - | - | - | ♦ | x | - | - | - | - | - | - | - | - | x | x | - | - |
| Sarkhy et al | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - |
| Ljungman and Myrdal | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - |
| Taghdir et al | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - |
| Wagner et al | - | - | - | - | - | - | - | x | - | - | - | - | - | - | - | - | - | - | - |
| Charalampopoulos et al | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - |
| Roma et al | - | - | - | - | - | - | - | - | - | - | - | x | x | - | - | - | - | - | - |
| Khurana et al | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - |
| MacCulloch and Rashid | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - | - |
| Garg and Gupta | - | - | - | - | - | - | - | x | - | - | - | - | - | - | - | - | - | - | - |
### Table 4 (Continued)

| Parent Related | Parent Related |
|----------------|----------------|
| Parents easily discuss child's condition | Parents felt child similar to other children |
| Parents easily interact with other parents | Parental knowledge of CD |
| Tendency to seek medical help | Concern about health of the child with CD |
| Parents consider child's diet to be a burden | Parents worry less about health |
| Parents consider child's diet to strain family budget | Concerned about future army service |
| Combined severity concerned | Evaluated parental knowledge of CD |
| Psychological burden on parents | Do not interact with parents of other CD children |
| Perception of the diet as a family burden |

| Authors | 
|---------|
| Mager et al | 
| Chauhan et al | x x x |
| Anon et al | 
| Sarkhy et al | 
| Ljungman and Myrdal | 
| Taghdir et al | 
| Wagner et al | 
| Charalampopoulos et al | 
| Roma et al | 
| Khurana et al | 
| MacCulloch and Rashid | 
| Garg and Gupta | ♦ ♦ |

**Abbreviations:** -, not studied; x, not found to be associated with compliance; ♦, found to be associated with compliance; ◊, associated with noncompliance; 1, established as a predictor after a logistic regression; 0, not established as a predictor after logistic regression; CD, celiac disease; GFD, gluten free diet; GI, gastrointestinal symptoms, ATIG.
### Table 5 Discrimination of variables regarding dietary, disease, personality, and quality of life-related factors

| Dietary related | Ability to choose GFD food | Ability to choose GFD beverages in menu | Reports child demands food not GFD once a week | Difficulty maintaining GFD in parties and special occasions | Shared responsibility in maintaining GFD | Availability of GFD | Hospital supply of GFD | Pricing of GFD | Budget spending on GFD | Perceived facility in maintaining diet | Difficulty to maintaining GFD at school | Difficulty to maintaining GFD at parties/ marriages | Likes gluten taste | No mistake in handling menu | Child perception of difficulty in maintaining GFD on travel | Child shares responsibility about disease |
|----------------|---------------------------|----------------------------------------|-----------------------------------------------|----------------------------------------------------------|------------------------------------------|-------------------|-----------------------|---------------|-------------------------|--------------------------------------|----------------------------------------|------------------------------------------|-----------------|---------------------|-----------------------------|-------------------------------------|
| Mager et al    | ✦                         | ✦                                      | ✦                                             | ✦                                                        | ✦                                        | ✦                 | ✦                     | ✦                         | ✦                       | ✦                                                                   | ✦                                                                     | ✦                                                         | ✦                             | ✦                     | ✦                     |
| Chauhan et al  | ✦                         | ✦                                      | ✦                                             | ✦                                                        | ✦                                        | ✦                 | ✦                     | ✦                         | ✦                       | ✦                                                                   | ✦                                                                     | ✦                                                         | ✦                             | ✦                     | ✦                     |
| Anson et al    | ✦                         | ✦                                      | ✦                                             | ✦                                                        | ✦                                        | ✦                 | ✦                     | ✦                         | ✦                       | ✦                                                                   | ✦                                                                     | ✦                                                         | ✦                             | ✦                     | ✦                     |
| Sarkhy et al   | ✦                         | ✦                                      | ✦                                             | ✦                                                        | ✦                                        | ✦                 | ✦                     | ✦                         | ✦                       | ✦                                                                   | ✦                                                                     | ✦                                                         | ✦                             | ✦                     | ✦                     |
| Ljungman and Myrdal | ✦                       | ✦                                      | ✦                                             | ✦                                                        | ✦                                        | ✦                 | ✦                     | ✦                         | ✦                       | ✦                                                                   | ✦                                                                     | ✦                                                         | ✦                             | ✦                     | ✦                     |
| Taghdir et al  | ✦                         | ✦                                      | ✦                                             | ✦                                                        | ✦                                        | ✦                 | ✦                     | ✦                         | ✦                       | ✦                                                                   | ✦                                                                     | ✦                                                         | ✦                             | ✦                     | ✦                     |
| Wagner et al   | ✦                         | ✦                                      | ✦                                             | ✦                                                        | ✦                                        | ✦                 | ✦                     | ✦                         | ✦                       | ✦                                                                   | ✦                                                                     | ✦                                                         | ✦                             | ✦                     | ✦                     |
| Charalam- popoulos et al | ✦                     | ✦                                      | ✦                                             | ✦                                                        | ✦                                        | ✦                 | ✦                     | ✦                         | ✦                       | ✦                                                                   | ✦                                                                     | ✦                                                         | ✦                             | ✦                     | ✦                     |
| Roma et al     | ✦                         | ✦                                      | ✦                                             | ✦                                                        | ✦                                        | ✦                 | ✦                     | ✦                         | ✦                       | ✦                                                                   | ✦                                                                     | ✦                                                         | ✦                             | ✦                     | ✦                     |
| Khurana et al  | ✦                         | ✦                                      | ✦                                             | ✦                                                        | ✦                                        | ✦                 | ✦                     | ✦                         | ✦                       | ✦                                                                   | ✦                                                                     | ✦                                                         | ✦                             | ✦                     | ✦                     |
| Mac-Culloch and Rashid | ✦                        | ✦                                      | ✦                                             | ✦                                                        | ✦                                        | ✦                 | ✦                     | ✦                         | ✦                       | ✦                                                                   | ✦                                                                     | ✦                                                         | ✦                             | ✦                     | ✦                     |
| Garg and Gupta | ✦                         | ✦                                      | ✦                                             | ✦                                                        | ✦                                        | ✦                 | ✦                     | ✦                         | ✦                       | ✦                                                                   | ✦                                                                     | ✦                                                         | ✦                             | ✦                     | ✦                     |
Table 5 (Continued)

| Disease Related          | Child perception of difficulty in maintaining GFD with friends | Dining in restaurants | Barriers to compliance: availability, cost, smaller communities | Age at diagnosis | Time since diagnosis | Comorbidities | Mean duration of CD | Serum ATG levels | Gluten intake | No positive history of CD | Self-reported GI symptoms | Frequency of CD dx in the family | Delay at diagnosis | Symptomatic patient | Type of manifestation | Celiac society membership | Symptom at diagnosis | Specific knowledge about the disease | Age at presentation | Cognitive restructuring |
|--------------------------|---------------------------------------------------------------|------------------------|---------------------------------------------------------------|-----------------|---------------------|---------------|---------------------|-----------------|---------------|--------------------------|--------------------------|-----------------------------|---------------------|-----------------------|----------------------|------------------------|------------------------|--------------------------|---------------------|--------------------------|
| Mager et al              | -                                                              | -                      | -                                                             | -               | -                   | -             | -                   | -               | x             | -                        | x                        | -                          | -                   | -                     | -                    | -                      |
| Chauhan et al            | -                                                              | -                      | -                                                             | -               | -                   | -             | -                   | -               | -             | -                        | -                        | -                          | -                   | -                     | -                    | -                      |
| Amon et al               | -                                                              | -                      | -                                                             | -               | -                   | -             | -                   | -               | -             | -                        | -                        | -                          | -                   | -                     | -                    | -                      |
| Sarkhy et al             | -                                                              | -                      | -                                                             | -               | -                   | -             | -                   | -               | -             | -                        | -                        | -                          | -                   | -                     | -                    | -                      |
| Ljungman and Myrdal      | -                                                              | -                      | -                                                             | -               | -                   | -             | -                   | -               | -             | -                        | -                        | -                          | -                   | -                     | -                    | -                      |
| Taghdir et al            | -                                                              | -                      | -                                                             | -               | -                   | -             | -                   | -               | -             | -                        | -                        | -                          | -                   | -                     | -                    | -                      |
| Wagner et al             | -                                                              | -                      | -                                                             | -               | -                   | -             | -                   | -               | -             | -                        | -                        | -                          | -                   | -                     | -                    | -                      |
| Charalamopoulos et al    | -                                                              | -                      | -                                                             | -               | -                   | -             | -                   | -               | -             | -                        | -                        | -                          | -                   | -                     | -                    | -                      |
| Roma et al               | -                                                              | -                      | -                                                             | -               | -                   | -             | -                   | -               | -             | -                        | -                        | -                          | -                   | -                     | -                    | -                      |
| Khurana et al            | -                                                              | -                      | -                                                             | -               | -                   | -             | -                   | -               | -             | -                        | -                        | -                          | -                   | -                     | -                    | -                      |
| MacCulloch and Rashid    | -                                                              | -                      | -                                                             | -               | -                   | -             | -                   | -               | -             | -                        | -                        | -                          | -                   | -                     | -                    | -                      |
| Garg and Gupta           | -                                                              | -                      | -                                                             | -               | -                   | -             | -                   | -               | -             | -                        | -                        | -                          | -                   | -                     | -                    | -                      |
| Personality Related | Quality of Life |
|---------------------|----------------|
| Problem-solving     | Higher HRQOL in physical domains |
| Social withdrawal   | Higher HRQOL in social domains |
| Wishful thinking    | QoL perceived by children (p = 0.0045) |
| Resignation         | QoL (diet parameter) evaluated by parents |
| Self-blame          | CDDUX score |
| Less emotional      | Parents CDDUX |
| regulation          |                |
| Less distraction    |                |
| Less blaming        |                |
| Novelty-seeking     |                |
| others              |                |

Abbreviations: -, not studied; x, not found to be associated with compliance; ♦, found to be associated with compliance; 0, associated with noncompliance; 1, established as a predictor after a logistic regression; 0, not established as a predictor after logistic regression; CD, celiac disease; GFD, gluten-free diet; HRQOL, health related quality of life; QoL, quality of life.
clinician interviews. Studies that used questionnaires had designed questionnaires specifically for the study or relied on previous validated questionnaires. Only one article used biopsies in follow-up. Despite doing so, the results were neither described nor used in conclusions. No article reported the use of urine gluten immunogenic proteins in monitoring disease activity. — Table 6 summarizes these findings.

### Age
The works evaluating age treat this variable as categorical, dichotomous, and use different cut offs. Independently of the cut off value, these studies usually consider two groups, “younger” and “older” children.

In five papers (three of “Good” quality, one of “Medium” quality, and one of “Poor” quality), younger age was significantly associated with compliance. In addition, two of these articles performed impact analysis, using logistic regression, and showed “older age” contributes to predict noncompliance. This observation was further supported by another two studies (one of “Good” quality and one of “Medium” quality) showing older children to be significantly associated with noncompliance. In our sample, two papers (of “Medium” quality and of “Poor” quality) were unable to report significant differences.

### Maternal Education
In our review, eight articles evaluated maternal education. Four articles (two of “Good” quality and two of “Medium” quality) showed maternal education to be a significant factor for compliance to a GFD while another four (two of “Good” quality and two of “Poor” quality) failed to show an effect.

Interestingly, one of the articles reporting an association between maternal education and compliance to a GFD, after performing a binary multivariate logistic regression analysis found this factor not to be a predictor of GFD compliance. In agreement, another article (“Good” quality) showed a correlation between lower maternal education and noncompliance to GFD.

### Paternal Education
Of the 12 articles included, six analyzed the influence of paternal education on a child’s compliance. Of these, two articles (both of “Medium” quality) showed paternal education to be a significant factor for compliance to a GFD whereas another four (two of “Good” quality and two of “Poor” quality) failed to do so.

### Parental Knowledge of CD
One article (“Good” quality) demonstrated this parameter as positively influencing compliance to GFD while another (“Medium” quality) found it not to influence it. Interestingly, one report (“Good” quality) distinguishes parental knowledge in perceived and evaluated, showing that only the first was significantly associated with compliance to GFD. Moreover, after performing a multiple logistic regression they showed it is a predictor of compliance. At last, an article (“Good” quality) reported statistical differences between the compliant and noncompliant groups, concerning parental knowledge of CD but after conducting a binary multivariate logistic regression analysis, demonstrated this factor was not a predictor of compliance to GFD.

### Family Type
Four articles evaluated “family type” as a dichotomous variable: a nuclear family—in which only the parents and their children live together, or a joint family—where the household inhabitants includes other family members. Two papers (“Good” quality) reported that belonging to a nuclear family increases GFD compliance. In addition, one of the articles after performing a multivariable logistic regression analysis showed it to be a predictor of GFD compliance. The

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**Table 6** Methods used to access compliance across the study

| Authors                      | Questionnaire | Biopsy | Clinical Evaluation | Anti-TG Antibodies | EMA | Antireticulum Antibodies |
|------------------------------|---------------|--------|---------------------|--------------------|-----|-------------------------|
| Mager et al                  | Applied       |        |                     |                    |     |                         |
| Chauhan et al                | Applied       |        |                     |                    |     |                         |
| Anson et al                  | Applied       |        |                     |                    |     |                         |
| Sarkhy et al                 | Applied       |        |                     |                    |     |                         |
| Ljungman and Myrdal          | Applied       |        |                     |                    |     |                         |
| Taghdir et al                | Applied       |        |                     |                    |     |                         |
| Wagner et al                 | Applied       |        |                     |                    |     |                         |
| Charalampopoulos et al       | Applied       |        |                     |                    |     |                         |
| Roma et al                   | Applied       |        |                     |                    |     |                         |
| Khurana et al                | Applied       |        |                     |                    |     |                         |
| MacCulloch and Rashid        | Applied       |        |                     |                    |     |                         |
| Garg and Gupta               | Applied       |        |                     |                    |     |                         |

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References:
1. Studies that used questionnaires had designed questionnaires specifically for the study or relied on previous validated questionnaires. Only one article used biopsies in follow-up. Despite doing so, the results were neither described nor used in conclusions. No article reported the use of urine gluten immunogenic proteins in monitoring disease activity.
2. Table 6 summarizes these findings.
3. The works evaluating age treat this variable as categorical, dichotomous, and use different cut offs. Independently of the cut off value, these studies usually consider two groups, “younger” and “older” children.
4. In five papers (three of “Good” quality, one of “Medium” quality, and one of “Poor” quality), younger age was significantly associated with compliance. In addition, two of these articles performed impact analysis, using logistic regression, and showed “older age” contributes to predict noncompliance. This observation was further supported by another two studies (one of “Good” quality and one of “Medium” quality) showing older children to be significantly associated with noncompliance. In our sample, two papers (of “Medium” quality and of “Poor” quality) were unable to report significant differences.
5. Maternal Education
6. In our review, eight articles evaluated maternal education. Four articles (two of “Good” quality and two of “Medium” quality) showed maternal education to be a significant factor for compliance to a GFD while another four (two of “Good” quality and two of “Poor” quality) failed to show an effect.
7. Interestingly, one of the articles reporting an association between maternal education and compliance to a GFD, after performing a binary multivariate logistic regression analysis found this factor not to be a predictor of GFD compliance. In agreement, another article (“Good” quality) showed a correlation between lower maternal education and noncompliance to GFD.
8. Paternal Education
9. Of the 12 articles included, six analyzed the influence of paternal education on a child’s compliance. Of these, two articles (both of “Medium” quality) showed paternal education to be a significant factor for compliance to a GFD whereas another four (two of “Good” quality and two of “Poor” quality) failed to do so.
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11. One article (“Good” quality) demonstrated this parameter as positively influencing compliance to GFD while another (“Medium” quality) found it not to influence it. Interestingly, one report (“Good” quality) distinguishes parental knowledge in perceived and evaluated, showing that only the first was significantly associated with compliance to GFD. Moreover, after performing a multiple logistic regression they showed it is a predictor of compliance. At last, an article (“Good” quality) reported statistical differences between the compliant and noncompliant groups, concerning parental knowledge of CD but after conducting a binary multivariate logistic regression analysis, demonstrated this factor was not a predictor of compliance to GFD.
12. Family Type
13. Four articles evaluated “family type” as a dichotomous variable: a nuclear family—in which only the parents and their children live together, or a joint family—where the household inhabitants includes other family members. Two papers (“Good” quality) reported that belonging to a nuclear family increases GFD compliance. In addition, one of the articles after performing a multivariable logistic regression analysis showed it to be a predictor of GFD compliance.
remaining two papers (“Medium” and “Poor” quality) showed no differences between compliant and noncompliant groups in relation to “family type.”

**Child’s Knowledge of CD**

The two works (of “Poor” quality) that analyzed this variable showed a child’s knowledge of CD as positively influencing compliance to GFD.

**Member of Celiac Society**

Two articles (one of “Medium” and one of “Poor” quality) evaluating the association between “adherence” and “celiac society membership” found significant differences between the compliant and noncompliant groups, with membership increasing GFD compliance. However, in one article, after performing a multivariate logistic regression analysis the effect was lost.

**Quality of Life**

Three articles were found concerning the QoL. One article (of “Good” quality) evaluated QoL on four different domains (social, emotional, school, and physical) and found that higher compliance was related with higher parental, perceived QoL, in social domains, and with child perceived QoL, in physical domains. Another article (of “Poor” quality) evaluated QoL based on another scale—socio-demographic, QoL, diet, communication, and having CD. Using this scale, only the “diet” section as positively influenced compliance to GFD. Using the same scale, the last article (of “Medium” quality) was unable to find any differences in the overall QoL.

**Maintaining GFD**

Three papers evaluated the difficulty in maintaining GFD. Two (of “Good” quality) stated children with higher compliance to the GFD considered gluten to have a “good taste” and found it less difficult to keep GFD generally at school and parties/marriages. Moreover, a binary multivariate analysis showed these factors were positive predictors of compliance. Another article (of “Medium” quality) found that children who reported more difficulty in maintaining GFD were more frequent on the noncompliant group.

**Discussion**

Compliance within pediatric age is challenging, hence the importance of finding factors associated with compliance. Overall, our review showed three main factors associated with GFD compliance, “Family type” and “Parental knowledge of CD” positively increase GFD compliance while increasing “age” in pediatric population decreases compliance. In addition, maternal education and Celiac Society Membership are also related with GFD compliance.

In comparison to other topics regarding CD, little literature can be found regarding this subject, especially in pediatric populations. Our first literature search was unable to identify any systematic work on this matter and the first review published on the topic was on June 4, 2019 by Myléus et al. In this work, Myléus et al reviews the methods used to evaluate GFD compliance and associated risk factors while we propose to identify specific factors that positively or negatively affect GFD compliance.

In addition, we found the need to address compliance specifically in CD patients without comorbidities since some of these, such as diabetes mellitus, imply additional lifestyle changes and diet restraints.

While it is logical to understand “age” as having a major role in compliance, our results confirm this parameter is a good predictor of GFD compliance. As age increases so does autonomy in food selection. Furthermore, with age, there is also the need for social integration and peer approval. Also, considering an early age of diagnosis, it is less likely for a patient to develop unrestricted eating habits. All these are challenging when maintaining a different lifestyle, such as a GFD. Despite the absence of a cut-off value, there is a general consensus among the literature as per which changes in compliance are associated with age. Additionally, two articles demonstrated “older pediatric age” was associated with prediction of noncompliance. In fact, they stated that with each year of increase in age, the child had a 25% less chance of remaining compliant to a GFD. Although one work reported opposite findings, its “Poor” quality limits its relevance to our analysis.

When analyzing children’s behavior toward maintaining a GFD, we expected to find differences between compliant and noncompliant children. Indeed, children that considered “gluten products as having better taste” displayed better GFD compliance and those reporting GFD “to be more difficult to maintain” showed less GFD compliance. While being quite challenging, as our data shows compliance decreases as children grow up, it is nonetheless a window of opportunity for pediatricians and dieticians to intervene by increasing the children’s knowledge of CD and, consequently, GFD may become a more achievable goal. It should be noted that as the ability to “maintain a GFD” was self-reported, the relation of causality might be misleading so further investigation is required.

Concerning the effect of maternal education, we would expect this parameter to be a good predictor of GFD compliance due to the mother’s role in food preparation and nutritional care. However, the results remain controversial as half of the studies found this parameter to be related to GFD compliance while the remaining did not, highlighting the need for further investigation.

To have a clearer picture on the influence of parental roles and since most works only study the maternal contribution, when available we also analyzed the influence of the father’s education to GFD compliance. Two “Good” quality studies with large populations were unable to demonstrate differences in the impact of this parameter between complaint and noncompliant groups, although smaller studies of “Medium” quality supported this hypothesis. According to our results, it is likely this factor is not associated with GFD compliance, but it should be further explored in future works.

It is important to stress that there is no evidence of correlating parental knowledge of CD with formal education. Nonetheless, parental knowledge of CD is related with compliance to GFD in pediatric populations, probably due to the major
role that parents play in choosing their child’s diet. In fact, Charalampopoulos et al showed parental knowledge is a predictor of compliance as children whose parents had a high perceived knowledge on CD were 3.3 times more likely to follow a strict diet. While this data was also supported by another “Good” quality work, two others of identical quality failed to identify it as a predictor of compliance emphasizing the need for further studies on this subject. Even though generalization is not possible at this time, physicians could still use parental knowledge of CD as a line of action to increase GFD compliance. This seems logical since parents with more knowledge will fail less in making diet choices, therefore their child complies more with a GFD.

Children feeding habits are the result of parental ability to drive their choices. Also, we report that decrease in compliance relates with older age. Therefore, it would be worth exploring this shift in their routine, as it comprises a change in compliance.

The analysis of “family” as a unit shows nuclear families are associated with compliance to GFD while joint families are related with noncompliance. One work demonstrated children from nuclear families to be four times more likely to maintain a GFD—an effect thought to be associated with parents being more focused on their child’s routine and environment. In a joint family, the amount of people eating on other diets may tempt the child to not comply. Nevertheless, two articles were unable to find a relationship between GFD compliance with the family type. Given the lower quality of this work, there is a limitation to the relevance of these findings in our analysis. Importantly, these findings concerned a geographically limited region limiting its generalization due to potential sociodemographic differences.

One would expect children to be more knowledgeable of CD to be more inclined to be GFD compliant given their raised awareness of the consequences of nonadherence and familiarity with gluten-free products. Indeed, two studies showed children’s knowledge of CD to positively influence GFD compliance, however, due to the small population size evaluated, the overall quality of these reports is poor and the vague definition of “knowledge in CD” extrapolation is limited.

Celiac Society Membership per se is positively related with compliance, which in light of our data are probably due to increased availability of information and contact with other CD patients. Importantly, the extent of its importance decreases with increasing levels of parental knowledge and with increased age of pediatric patients. It is important to note that only two studies demonstrated this factor as influencing compliance, one of them having “Poor” quality. Also, these results were based on specific populations (Greek), and therefore a selection bias may be present which partially limits its extrapolation to other populations.

High heterogeneity is observed regarding QoL. This is due to the high variability of tools used to assess it and to the inconsistency of the studies overall quality. Therefore, it is not possible to draw conclusions concerning this parameter.

Several factors limited our analysis. First, the heterogeneity of the tools used to measure GFD compliance leads to inconsistent results between reports. Standardization of research protocols would greatly enhance the quality and validity of future studies. Also, without having an objective measurement, such as t-TG levels, it is difficult to exclude contaminations.

Second, the definition of “GFD compliance” needs to be clarified. In the studies herein included, compliance was broadly defined as “a dichotomous variable in which a positive outcome is identifiable as the absence of awareness of gluten intake.” By contrast, any awareness level of transgression to a GFD was considered as noncompliance. Our findings are corroborated by Myléus et al since they also could not find any method more reliable to assess compliance.

Third, most studies rely on parental information alone; which by itself is a potential source of bias since patients themselves do not report data. Of course, given the specific particularities of this population, it might not have been possible to obtain the data otherwise.

Fourthly, an additional source of bias was associated with the lack of standardization in the age categories at study. The lower and higher limits of each age group varied greatly between reports, compromising its evaluation. Finally, there was an overall lack of quality in what concerns study design and the presentation of results.

As to the reports’ quality assessment and subsequent extrapolation of results, we would like to note that since the majority of the studies were cross-sectional, observational by nature, the results obtained could not be standardized as predictors, as the relationship between exposure and outcome cannot be fully discriminated. For this same reason, in the quality assessment, most studies consistently received a negative score for the study design. Additionally, we stress the need for further studies that establish a correlation between exposure and outcome.

**Conclusion**

Undoubtedly, GFD is strongly related with social and environmental contexts. The interconnection between parental features and children outcomes became clearer. Despite some of them being unchangeable, such as age or family type, with this work, we were able to show evidence that modifiable factors, parental knowledge on CD per example, can also play major roles in compliance. Nevertheless, the causality between these factors and compliance still remains unclear. Therefore, the need for further knowledge in causality relations is in order so that compliance rates, among pediatric populations, can be increased.

Regarding this, age was proved to influence compliance as young children consistently display better compliance rates. More so, children of informed parents and a nuclear family household were independently positive influencers of compliance. All these predictors should be taken into account in clinical practice when evaluating CD patients.

Lastly, it is important to state that a systematic approach to compliance should be established, only then GFD compliance will be fully understood.
Authors’ Contributions
V.M.-C. and R.M.-C. conceptualized the work, collected, analyzed, and interpreted, drew and reviewed the first draft and ensured for the integrity and accuracy in the work. H.A. conceptualized the work, reviewed the first draft, and ensured for the integrity and accuracy in the work. F.P.-R. reviewed the first draft and ensured for the integrity and accuracy in the work. All the authors approved the final manuscript as submitted and agree to be accountable for all aspects of the work. These authors contributed equally to this work.

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Conflict of Interest
None declared.

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