Abstract: Gluten is a complex mixture of storage proteins in cereals like wheat, barley and rye. Prolamins are the main components of gluten. Their high content in proline and glutamine makes them water-insoluble and difficult to digest in the gastrointestinal tract. Partial digestion generates peptide sequences which trigger immune responses in celiac and gluten-sensitive patients. Gluten detection in food is challenging because of the diversity, in various food matrices, of protein proportions and their immunogenicity. Attempts to develop standard reference materials have been unsuccessful. We present here a summary of recent studies reporting the detection of dominant Gluten Immunogenic Peptides (GIP) sharing epitopes presented in the α-gliadin 33-mer, the most important celiac disease-immunogenic sequence within gluten. GIP were not only detectable and quantifiable in very different kind of difficult to analyze food, but also in stool and urine of celiac patients on a supposedly gluten-free diet (GFD), providing the first simple and objective means to assess adherence to the GFD. Methods to specifically and sensitively detect the most active GIP in food and biological fluids are rational candidates may use similar analytical standard references for determination of the immunopathological risk of gluten exposure in gluten-related diseases.

Keywords: gluten immunogenic peptides; celiac disease; gluten quantitation; gluten food analysis;

1. Introduction

Gluten is a complex mixture of storage proteins found in cereal seeds composed of hundreds of related but distinct proteins, mainly prolamins and glutenins [1]. Wheat gluten proteins have been classically defined according to their solubility [2]. In wheat, they are categorized as albumins (soluble in water), globulins (soluble in saline solutions), gliadins (soluble in 60-70% ethanol) and glutenins (only soluble under stronger conditions, i.e. acids, reducing agents and detergents, urea, etc) [3]. According to their electrophoretic mobility, gliadins are subdivided into three main
groups: the first α- and β-gliadins, the second Υ-gliadins and the third consists of Ω-gliadins [2].

Prolamins contribute to the cohesiveness and extensibility of the gluten, whereas glutenins play a role in the maintenance of the elasticity and strength of the gluten [4]. Prolamins have been widely studied due to their contribution to the quality of the end product of bakery and pasta foods, including the rheological characteristics of dough made from wheat flour [2,5]. Prolamins are the major storage proteins in wheat (gliadin), barley (hordein), rye (secalin), corn (zein), sorghum (kafrin) and a minor protein in oats (avenin) and rice (orzein) [6]. Oats species have the lowest variability in prolamin content; wheats have the highest content (Figure 1).

High densities of proline and glutamine residues are present in the prolamins which are synthesized and deposited in the endosperm of the grain as primary source of nitrogen for protein synthesis, which occurs later during germination [12–14]. The proteins and polypeptides within the Prolamin Superfamily possess similar structures: signal peptide for translocation into cellular compartments, a non-repetitive N-terminal region, a non-repetitive C-terminal region and a long repetitive central region. The central region contains glutamine rich and proline-rich repeat units unique to each group [14]. However, there are differences in the number, content and properties of prolamin polypeptides among these cereals. Strictly speaking, the term gluten could comprise all prolamins including those of rice or maize. However, for simplification and practical reasons, the term gluten is typically restricted for prolamins with the ability to trigger immunotoxicity in certain groups of patients with gluten-sensitive spectrum, which includes three main forms: allergic (wheat allergy), autoimmune (celiac disease, (CD), dermatitis herpetiformis and gluten ataxia), and immune-mediated (gluten-sensitivity). This simplification often leads to confusion when determining the gluten content in different cereals by different immunologic methods. To understand the factors affecting gluten content estimations, we evaluated the recent literature on methodologies for detecting gluten, as well as the reference material for the relevant articles. The aim of this review article is to provide an overview of the current definition of gluten according to the latest methods and scientific understanding, and to discuss the appropriate standards to assess immunogenic gluten in foods and human biological samples.

The search was conducted in PubMed MEDLINE and SCOPUS databases. The following search terms were used: “gluten content”, “gluten analysis”, “celiac immunogenic peptides”, “gluten immunogenic peptides” and “toxic gluten”. The keywords “toxicity and celiac disease”, “immunogenicity and celiac disease”, “harmful prolamin and celiac disease”, “gluten proteins and celiac disease”, “gluten immunogenic peptides and celiac disease” and “gluten peptides and celiac disease” were also used.
2. Gluten as the trigger for autoimmunity

Gluten proteins found in wheat, barley, and rye are among the only food ingredients known to trigger an autoimmune condition: celiac disease (CD) [15]. Exposure to gluten in combination to genetic predisposition and other unknown immunologic and environmental factors can trigger CD [16]. The main pathological lesion produced by gluten in CD is located in the proximal small intestine; however, the disease is increasingly considered a systemic disorder rather than a disease limited to the gastrointestinal tract. This is primarily explained by the fact that CD belongs to the group of autoimmune diseases (AD). In the full-blown lesion, there is loss of small bowel intestinal villi and infiltration of leukocytes, both in the epithelium and in the lamina propria. The chronic inflammatory condition of CD can be reversed with a gluten-free diet (GFD) [17]. The symptoms associated with gluten intake in CD can be diverse and include diarrhea, constipation, anemia, osteoporosis, dermatitis herpetiformis, fatigue, and infertility [18]. Moreover, a significantly increased prevalence of other AD has been reported in individuals with CD and their first-degree relatives as compared to controls, with an estimated burden of AD in CD cases up to 15% [19,20]. It has been suggested that these associations among CD and other AD may be explained by the sharing of a common pathogenic basis involving genetic susceptibility, similar environmental triggers, and the loss of intestinal barrier secondary to dysfunction of intercellular tight junctions with increased intestinal permeability, and possibly by other undiscovered mechanisms [19,20]. The type of symptoms and their intensity is largely dependent on the individual, and likely related to the pattern and amount of gluten consumption [21]. The HLA-DQ2 and HLA-DQ8 genotypes determine the risk of disease development because the corresponding encoded HLA-DQ2 and HLA-DQ8 bind to gluten peptides and present them to the T lymphocytes [22].

After ingestion of gluten-containing food, gluten is partially digested by enzymes of the gastrointestinal track into relatively large peptides [23]. Some of these peptides can bind directly to HLA-DQ2.5 or -DQ8 and trigger T-cell responses, which may result in local tissue damage [15,24]. In parallel, those gluten peptides can trigger the release of tissue transglutaminase 2 (TG2) which can deamidate some glutamine residues to glutamate in a large number of gluten peptides [15,25]. Deamidated gluten peptides can bind with much higher affinity to HLA-DQ2.5 or -DQ8, which can amplify the gluten-specific T-cell response. The activation of TG2 is thus a crucial step to increase the immunotoxicity of the gluten peptides [24]. The gluten-specific T-cell response induces the release of inflammatory cytokines that generate local inflammation [26].

3. Immune activating potential of gluten peptides

Hundreds of immunogenic peptides have been described in wheat, barley, rye and oats, which are able to activate T cells [11,23,27]. However, not all gluten peptides are equally harmful to CD patients, as the number of epitopes present in
these peptides may vary strongly between different cereals [28] and between cereal species [29]. Ciccocioppo et al. [30] suggested that gluten peptides are divided in two groups, toxic and immunogenic. Both pathways interact and potentiate each other to sustain the chronic process of the intestinal damage. Toxic peptides are capable of inducing non T-cell mediated mucosal damage when administered “ex vivo” on biopsies from celiac small intestine. Gluten Immunogenic Peptides (GIP) contain sequences which specifically stimulate T-cell lines isolated from peripheral blood of CD patients. Once a peptide is identified as immunogenic, it is considered relevant for those with CD, regardless whether or not it has also toxic properties [31].

The complete repertoire of peptides involved in the pathogenesis of CD remains a daunting task because of the great heterogeneity of gluten proteins [24,27,32–37]. Several studies have demonstrated that peptides derived from α-gliadins induce strong responses in the large majority of patients, while responses to the other peptides are less frequently found. It is possible that some immunogenic gluten epitopes may be tolerated by CD patients depending on the patients’ sensitivity to the different immunogenic epitopes [15,38,39]. In this scenario it becomes very important to accurately detect which immunogenic gluten epitopes are present in foods.

Several gliadin peptides have been described to induce the adaptive immune response, but most of them are digested by gastric, pancreatic and intestinal proteases. Two well-known main peptides remain undigested: the 33-mer (P55-87) and the 25-mer (P31-55). Consistently, these two peptides are active “in vivo” in the celiac intestine after gluten ingestion [40–43]. A peptide sequence located at position 31-43 of α-gliadin, which is contained in the 25-mer, represents a toxic peptide [44–46]. The P56-68 of α-gliadin, which is contained in the 33-mer, is considered to be the most important CD-immunogenic sequence within gluten [24]. The α-gliadin 33-mer (amino acid sequence LQLQPFPQPQLPYPQPQLPYPQPQLPYPQPQPF), contains three repetitions of the P56-68 of α-gliadin [43]. This peptide is present in the N-terminal repetitive region of α-gliadins and contains six overlapping copies of three different DQ2-restricted T-cell epitopes with highly stimulatory properties [43]. It harbours the p56-75 peptide (LQLQPFPQPQLPYPQPQPLPY) that has been identified as the dominant gluten epitope [38,39]. The presence of the 33-mer in most common wheat and spelt cultivars have been recently detected and quantified by LC-MS/MS (Liquid chromatography tandem-mass spectrometry) ranging a level of 91-603 μg/g flour [47]. The special focus in the literature on this most immunodominant peptide seems to be justified.

Whereas the pathogenic role of CD4-mediated cells in CD is well-defined, several other studies have pinpointed the involvement of an innate, non-T-cell mediated immune response in intestinal mucosal atrophy [48]. Interestingly, among these peptides, two sequences of 18-mer and 25-mer lengths were truncated versions of the well-described 33-mer, and one peptide corresponded to the innate immune eliciting peptide 31-49 [33].
It is not fully understood how GIP can pass the epithelial barrier into the lamina propria to induce immune responses. Studies using biopsies from control and celiac patients found that P57-68 was completely degraded after mucosal to serosal transport. In contrast to biopsies of celiac patients, P57-68 was already partially degraded by brush-border peptidases in control subjects [49,50]. However, the use of intestinal biopsies from celiac and control patients to investigate the transport and degradation of GIP involves a complex interplay between various cell types and hardly allows to reveal the role the enterocytes in the metabolic fate of GIP and thus their role in antigen transport and presentation. The transport in the gut epithelium of two gluten peptides and among them, the immunogenic P56-68 peptide as representative GIP, has been studied [51]. GIP transport and increased uptake depended on the chain length as well as the integrity of the epithelial barrier system [51]. The absorption of gliadin peptides is complex. Reports indicate that GIP may be uptaken by the paracellular route [52]. Bruun et al. [53] also demonstrated that gluten fragments cross the intestinal barrier to be distributed to organs other than the gut, finding these gluten peptides in pancreas by mass spectroscopy and hypothesized the initiation of inflammation and induction of beta cell stress. The detection of gluten peptides in urine by using anti-gliadin 33-mer monoclonal antibodies (moAbs) demonstrated that gluten peptides can be absorbed, transported in the blood stream and excreted [54,55].

A systematic study of hundreds of gluten epitopes by ELISPOT revealed that three highly immunogenic peptides, derived from α-gliadin (ELQPFQPPELPYPQPQ), ω-gliadin/C-hordein (EQPFPQPEQPFPWQP), and β-hordein (EPEQPIPEQPQPQ), account for 90% of the celiac-specific immuno-response elicited by the full proteome of wheat, barley and rye proteins [27,39,43,56]. Recently, a HLA-DQ gluten tetramer test comprising only five gluten T cells epitope sequences (Table 1) could serve to identify celiac patients with 100% sensitivity and 90% specificity vs. control with gluten containing diet [57]. Those results also suggested that a limited number of peptides represent more than 90% of the celiac specific T-cell response.

4. Analysis of gluten peptides

Many methods have been developed over the years for the detection of prolamins, including the polymerase chain reaction (PCR), LC-MS, and immunological methods based in antigluten peptide antibodies. Immunoassays as ELISAs (enzyme-linked immunosorbent assays) and LFDs (lateral flow devices) have been the method of choice because of combination of specificity, sensitivity, simplicity and cost effectiveness in the food industry to certify gluten-free food. An ideal Ab for gluten analysis should be not only a reliable indicator of the presence of prolamins from cereal species known to be toxic to CD patients but also should recognize the specific intramolecular regions responsible for such toxicity and immunogenicity [1].
The lack of an official reference material; the diversity of matrices, target analytes, sample extraction buffers, extraction time and temperature, as well as calibration standards; and differences in Ab specificity are factors contributing to the differences between the various methods of detection and quantification of gluten [62-63].

Several antibodies (Ab) have been raised against different prolamin epitopes (Table 2). Skerrit and Hill [64,65] developed a sandwich format that was approved as an official method by AOAC (Association of Official Agricultural Chemist) and it was used for many years in gluten analysis [1]. Other antibodies were raised against different epitopes of α-gliadin, such as PN3 (residues 31-49) for the toxic 19-mer peptides [45,66], CD5 (residues 51-75) or Abs against T-cell stimulatory peptides present in gluten [67]. The R5 moAb, raised against Ω-secalin from rye [68-69] recognizes highly repeated peptide sequences present in wheat, barley and rye grains. R5 reacts with high sensitivity against the epitope QQFP as well as QLPFP, LQPFP, and QQQFP present in celiac-toxic sequences that occur repeatedly in gliadins, hordeins, and secalins [68]. However, when the reactivity to those epitopes has been brought to experimental testing, the sensitivity for some of the main immunogenic peptides, such as the gliadin 33-mer, was not significant [61,70-71]. The poor recognition of some of the main immunodominant gluten peptides may underestimate the immunogenicity of some hydrolyzed food and beverages [61].

Two moAbs, G12 and A1, were raised by the authors against the main immunogenic epitope of the α-gliadin 33-mer. These antibodies were able to detect the presence of gliadin 33-mer-related epitopes in prolamins from wheat, barley, rye and various oats varieties [58,72,75], as well as in food samples [60-61,76] and human samples to monitor GFD compliance and transgressions [54,59,77-78]. This technique has served to assess the efficacy of new experimental drugs or strategies to eliminate GIP during digestion [43,72,75,79].

Another anti 33-mer Abs, 14G11 and 13F6, have been recently used for the portable sensor device, Nima Sensor [71]. In addition to the affinity for the immunodominant gliadin 33-mer peptide not detected by R5, 14G11 and 13F6 affinity for gliadin also was 35 and 6.6 fold higher than the affinity of the R5, respectively. The performance of that device was tested in comparison with ELISA R5 and G12. Estimations in spiked samples with 10, 20 and 30 parts per million (ppm) of gluten in 13 different food matrices revealed that the device was able to detect 75.6%, 87.5% and 93.6% positives. Intriguingly, the Codex Alimentarius Type I ELISA Sandwich R5 was underestimating gluten content in 13 out of 13 food samples with intended 20 ppm of gluten content in that study and 7 out of 13 for ELISA G12 [80].

5. Factors affecting gluten content estimations

Despite the lack of standardized reference material for gluten, there is a Codex Alimentarius Type I official method based on the ELISA R5 and Mendez Cocktail
However, there are many food matrices difficult to measure due to their interferences with antibody binding, cross-reactivity, or problems in extracting the gluten [31]. In addition, there are open questions and wrong assumptions in assessing the gluten content:

5.1. Gliadin as standard

The most typical calibration standard is gliadin because it is from wheat, which is the most frequent source of gluten; it is at least soluble in ethanol-water assuming that the proportion gliadin/glutenin in gluten is 1/1. In the estimation, gluten content is calculated to be 2 times the estimated gliadin content. However, the proportion prolamin/glutenin varies greatly among the different cereals and cultivars [82]. The underestimation or overestimation due to that assumption may be more frequent than initially assumed.

5.2. Is gluten equivalent in wheat, barley, rye?

The reactivity and number of immunogenic peptide sequences may vary among different wheat [47,83-84] and barley [42] varietals. All wheat flours studied by Schalk et al [47] contained the 33-mer peptide. In contrast, the 33-mer was absent (<limit of detection) from tetra- and diploid species (durum wheat, emmer, einkorn), most likely because of the absence of the D-genome, which encodes α2-gliadins. In Comino and coworkers [42], eight different barley cultivars were analyzed by ELISA G12 which revealed 25-fold differences in reactivity between the most and the least reactive barley cultivar. Three of those cultivars were analyzed by T-cell activation, and the hierarchy of immunogenicity with T cells isolated from peripheral blood was consistent with the reactivity of the barley kernels. Therefore, the reactivity of the monoclonal antibodies used in the detection of gluten content may provide different estimations that should be verified with the real immunogenicity with human samples.

5.3. Can oats contain gluten?

Oats contains about 15% of avenin. However, this prolamin has not been broadly accepted to be named as gluten, in order to restrict the term ‘gluten’ to those prolamins generating clear food intolerance to certain individuals. It has been assumed by many stakeholders that oats are naturally gluten-free and only when oats are contaminated with other toxic cereals as wheat or barley, reactivity in gluten ELISA tests could be observed [85]. However, some pure oats cultivars have significant cross-reactivity with the most used monoclonal antibodies R5, G12 or Skerrit [58,86-87]. Some celiac T-cell activating sequences from oat have been identified [11,88-89] and some oat varieties have elicited early inflammatory events typical of CD [90]. Despite these evidences, it is still commonly believed that there is
no reactivity in pure oats. However, most of the clinical studies where gluten challenges were made with non-reactive oats-assessed with the most commonly used immunomethods- suggest that oats may be safe for most celiac patients [89,91]. It remains to be verified whether 2 out of 15 patients with persistent histological deterioration after one year on oats challenge are due to lapses in the GFD or to the oats peptides [91]. In any case, it seems that lack of reactivity with immune assays (R5, G12, Skerrit, etc) may guarantee the absence of gluten regardless of source - from oats itself or from wheat or barley contamination.

5.4. How much immunogenic gluten is there in hydrolyzed gluten?

Fragmented gluten proteins can be found in beers, baby food, etc. The potential diversity in the generation of sequences, relative abundance and extension of the resultant gluten peptides is almost limitless [92]. The estimation of gluten equivalence in hydrolyzed gluten samples is thus a challenge. Firstly, peptides may have only one epitope per molecule. Therefore, the sandwich ELISA R5 underestimates the gluten peptide content of some beers [93]. This problem was solved by using competitive R5 ELISA that could measure single epitope-peptide. However, semi quantitative analysis of one hundred Belgium beers by a sandwich based A1/G12 lateral flow strips showed consistent results compared to a competitive G12 ELISA [60]. The most reactive fractions in High Performance Liquid Chromatography (HPLC) analysis for the G12-based immunomethods of samples of those beers also provided the highest immunoactivity. The G12 most reactive fractions also allowed to identify immunogenic gluten peptide sequences by Matrix assisted laser desorption/ionization-time of flight (MALDI TOFF/TOFF). Further analysis of one of the beers with differences in estimated gluten content by competitive R5 and G12, revealed that many of the identified immunogenic peptides showed tandem epitopes for A1/G12, explaining the consistency of the results with sandwich and competitive immunoassays [76]. The difference in estimations between different antibody-based method could be more appreciated in hydrolyzed food or beverages because of differential resistance of the corresponding epitopes observed [94]. Furthermore, the resistance of the gluten immunogenic peptides with G12 epitopes allowed to detect gluten in fecal and urine samples [54,59]. The amount of gluten in biological fluids was estimated by using a gliadin 33-mer calibrator, enabling the estimation of the mean daily gluten consumption in celiac patients [55].

The capacity of G12 Ab to immunocapture most of the T-cell activity of either a barley beer or a pepsin-tripsinated gliadin was also tested by using a G12-agarose resin for immunoaffinity separation of G12 binding polypeptides [61]. Less than 5% of the polypeptides of the beer or the hydrolyzed gliadin were retained in the G12 column, however, the captured peptides were able to generate about 90% of the immunogenicity estimated by celiac T-cell activation. Interestingly, 25% of the beer polypeptides and 65% of the gliadin hydrolysate that were able to be detected by R5 Ab were also retained in the G12 immunoaffinity resins. Those results indicated
that most of the immunogenicity are detected by the G12 moAb and that the most used R5 Ab may underestimate the potential immunogenicity of certain hydrolytic material (Figure 2).

5.5. How much gluten does genetically-modified low immunogenic cereal contain?

Biotechnological approaches, such as gene silencing by RNA interference (RNAi), have been used to produce low-gluten wheat in recent years. It was found that the silencing of γ-gliadin genes affects the rheological properties of wheat dough, where the downregulation resulted in dough that were stronger and more tolerant to overmixing [95-96]. As indicated in Table 3, many of the immunogenic gliadins were suppressed one to two orders of magnitude, with regards to the wild types, in a group of six modified plants by iRNA [97]. The reactivity for R5 epitopes decreased to less than 2% in some lines with regards to those of the wild types (from 1.227 ppm that of the transgenic plant to 64.395 ppm -114.043 ppm).

However, the glutenin content of such transgenic plants did not decrease significantly (75% to 143% of HMW glutenin with regards to the wild types), and neither did the protein level (13.1 to 15.7 %). T-cell activation was tested with gluten extracts of the six transgenic plants. There was poor or no significant T-cell response in those six transgenic plants. Although immunogenic peptides have been described in HMW or LMW glutenins [23], those results with transgenic plants indicated that the immunoactivity of glutenins could be one order of magnitude lower than those of the gliadins.

Recently, Sánchez-León et al. [98] have used the CRISPR/Cas9 technology to precisely and efficiently reduce the amount of α-gliadins in the seed kernel, providing bread and durum wheat lines with reduced immunoreactivity for gluten-intolerant consumers. ELISA tests with R5 and G12 moAbs showed a strong reduction in gluten content (up to 85%) in the modified lines compared to that of the wild type.

The first “ultra-low-gluten barley” was produced in Australia [99] using traditional breeding techniques, where three recessive alleles were combined to create low-hordein parental genotypes. The gluten content of barley variety was reduced to below 5 ppm estimated by ELISA. However, in that study the gluten content was not determined with an antibody method that recognized GIP.

5.6. Gluten immunogenic peptides modified by bacterial enzymes

The use of protease enzymes to inactivate gluten peptides has also been investigated as a method to remove the CD antigenicity of wheat and other cereals. The challenge lies in preventing the degradation of the protease enzymes in the gastrointestinal tract [100]. A pool of selected lactobacilli demonstrated the capacity of strongly hydrolyzed the wheat bread gluten (18,000 ppm) to less than 10 ppm after 360 min of treatment by R5 sandwich and competitive ELISA [101]. The specificity of G12 moAb to detect GIP was exploited to evaluate the immunogenic potential of the pool of peptides produced during bacterial colonization [102]. Wei
et al. [94] found that exceptionally high gluten-degrading enzyme activities belonged to subtilisin protease family could eliminate of the major gliadin epitope in preventing T-cell activation in the lamina propria and G12 ELISA assay was employed to monitor the epitope elimination.

On the contrary, Janssen et al. [103] investigated the effectiveness of existing digestive enzyme supplements claimed to aid in gluten degradation by the R5 ELISA. Despite the observed reduction in the gluten content with the enzyme supplements by R5 method, authors argued that it did not reflect an actual breakdown of all immunogenic sequences because of being not enough specific for immunodominant peptide sequences from α-gliadins but is present in a number of γ-gliadin sequences. They corroborated by mass spectrometric analyses the gluten degradation products and certainly, the commercial enzyme supplements were completely ineffective in degrading immunogenic gluten fragments from both the α- and γ-gliadins.

6. Conclusions and Future Directions

After decades of gluten analysis by immune methods, there is no consensus reference material for gluten. The definition of gluten is still an unsolved issue. The amount of cereal varieties, proteins, peptide sequences, hydrolytic and other industrial processes, as well as the diverse individual response to gluten peptides, make the detailed analysis of the potential risk of food stuff extremely complex. However, it is reasonable to propose that the concept should be linked to the immunogenic activity in celiac patients. In the last years, the advance in the scientific knowledge and pathology of CD has allowed to identify not only gluten peptide sequences that trigger the immunological response by T-cell activations in celiac patients, but also the most immunodominant among them. There are already immune methods with reported characterization of the consistency with “ex vivo” quantified immunogenicity with T-cells from celiac individuals. A limited set of dominant GIP has already been characterized as responsive of most T-cell response in CD. GIP could be detectable in very different environments including hydrolyzed food and beverages, excreted in stool and urine, what is consistent with the resistance to digestion of GIP and the generation of systemic symptoms of gluten intolerant populations. A limited number of GIP could be obtained by synthesis at high purity or quantity, and could be used as calibrator of either immune methods or mass spectrometric analysis. We propose that they should be considered as future reference material to assess immunogenic gluten in food stuff.
**Patents:** The following patents are part of this work: Determination of levels of immunogenic gluten peptides in human samples (WO2012089868) and Detecting gluten peptides in human fluids (WO2016005643).

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