Celiac disease – Using science to develop a rational approach for screening of newly expressed proteins

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Searches with 8 aa peptides to identify allergens have been deemed to have little value.

Celiac specific peptide science is in a much better place:

- Organisms that cause the celiac disease are known
- Numerous peptides that bind cellular receptors and initiate the celiac amplification cascade are known

Regulatory guidance should leverage **all available scientific knowledge** to identify proteins with putative safety concerns.

Specificity that drives **consistency in the application of the guidance** is essential.
- Identify a comprehensive set of wheat, barley, rye and oat peptides that trigger celiac enteropathy
- Model the comprehensive set of peptides to develop rules
- Modify the guidance to drive a consistent approach to the following:
  - Thorough informatics screening of any food protein for possible similarity with hazardous celiac proteins
  - A step-wise process that is unambiguous
  - Start the hazard identification with the knowledge of the protein
  - Conservative approach that can identify potentially hazardous proteins/peptides
  - An informatics approach that is accurate, high-throughput, and can be interpreted by all stakeholders
For such screening to be useful it must display a reasonable level of specificity:

- **Example**: ELISA analyses are validated to show specificity for a target protein in a matrix – if not specific, they are useless in a complex matrix.

- In bioinformatics, the large databases are our “complex matrix” and our searches must be relatively specific to find relevant matches.

The draft guidance document proposed screening with a degenerate peptide sequence [EQ][LQFS]P[YFAQV]
Evaluation of Specificity by Searching SwissProt and Genbank

Specificity can be assessed by counting and determining the identity of aligning sequences

Imagine a bioinformatic screen that aims to identify all celiac peptides….

<table>
<thead>
<tr>
<th>Specificity</th>
<th>Proteins identified that are known to cause celiac</th>
<th>The rate of matches for proteins that don’t cause celiac</th>
</tr>
</thead>
<tbody>
<tr>
<td>High</td>
<td>&gt;90%</td>
<td>&lt;10% - good specificity</td>
</tr>
<tr>
<td>Medium</td>
<td>~50%</td>
<td>~50% - balanced specificity</td>
</tr>
<tr>
<td>Low</td>
<td>&lt;1%</td>
<td>&gt;99% - poor balance – cannot identify hazards because they are not separated from safe proteins</td>
</tr>
</tbody>
</table>
The Degenerate 4 Amino Acid Peptide Has Low Specificity – Inaccurately Identifies Safe Proteins in Common Foods

Survey of UniProt
SwissProt database
552,000 entries
(Esmeralda Posada, Bayer CropScience N.V.)

Survey of GenBank
49.5 million entries
(Andre Silvanovich, Monsanto company)

- 57,096 aligning proteins spanning 2,494 different species
- Within these species many are safely consumed by CD patients
  - Onion, garlic, cucumber, peanut, sugar beet, coffee
  - Beef, chicken, turkey,
  - Watermelon, strawberry, lemon orange and apple
  - Greater than 4,000 potato proteins
- Not possible to distinguish celiac peptides in a meaningful way with degenerate 4 aa peptide
- Modeling of the specific celiac peptides shows us a 4 aa peptide can be improved
- Celiac peptides are very specific, specificity occurs at positions outside of the 4 aa peptide

- 3.5 million produce an alignment
- Proteins from tens of thousands of species including humans, soy and corn were identified
- Among corn proteins were the high abundance zeins, a prolamin storage protein
Extending the Degenerate Peptide Length Decreases False Alignments while *Retaining Specificity*

Consensus amino acids

Proteins that require peptide modelling

Number of false positives is reduced

Using Science to Develop Screening of Novel Proteins for Celiac Hazard
Using a common denominator approach

- Shorter peptides are tested to determine if they are contained in larger peptides
- If a larger peptide contains a shorter peptide, the larger peptide is dropped
- If a protein contains a longer peptide, it will by definition also contain the shorter peptide, no need to use the long peptide as a query
- The 464 peptides are reduced to 160 by employing this filter step

```
PQQPFPQQ
PFSQQQQQ
PFPQPQLPY
PQPQLPYPQ
......
QQFLQ PQQPFPQQ PQQPYPQ
LQPQQPFPQQ PQQPYPQQP
QQQFIQPQQPFPQQ PQQTYP
```

This becomes the “screening peptide”

- Screening GenBank with the **160 peptides** yields 20653 exact matches
- Indicates an acceptable level of specificity from a screening perspective and is conservative enough to reliably identify celiac peptides
The Degenerate 4 Amino Acid Peptide

7 AA degenerate is the fullest form of celiac peptide screen

Better specificity here

Not specific enough – Degenerate 4 amino acid peptide yields 103 hits in the 160. Perhaps several degenerate peptides are needed?

3,530,146 (103/160)
502,837 (93/160)
139,000 (61/160)
64,659 (47/160)
Goal

- To identify a screening sequence of amino acids that describes the 9-mer peptides associated with Celiac disease

Premise

- There are 9-mers AND there are longer peptides that have been listed and associated with wheat, barely and rye that also contains these smaller (9 AA) peptides

Proposed Screening Objective

- Determine the smallest screening sequence or set of sequences that identify both the 9-mers and the longer sequences.
Proposed Informatics Workflow

Full length proteins in allergen database assessed for Codex Alimentarius criteria

Establish a refined “evergreen” list of short non-redundant Celiac Peptides

Use in process called “exact matching”

Establish screening algorithm with degenerate peptides

Use all known CD peptides and modelling to develop an “evergreen” set of degenerate peptides

Any significant matches between a novel protein and celiac peptide are removed from “development”, or considered for further modeling/in vitro screening

This is already performed in current processes
Guidance should take into account origin and knowledge on newly expressed proteins

Current draft guidance has a high false positive rate and is likely to miss CD proteins
  – False positives and risk of missing important matches limit the goal of conservative safety screening

A comprehensive list of celiac peptides should be evergreen and reside outside of the guidance document
  – Keeps the safety screening up to date

Any bioinformatics that includes degeneracy should be based upon modelling and the comprehensive evergreen list
  – Modelling supports the most accurate way to build a screening process – degeneracy alone is broad, but not accurate, so limits the goal of finding hazards

As we have seen with the 8-mer allergen search, old guidance does not go away
  – Limits the ability to deliver to the newest science