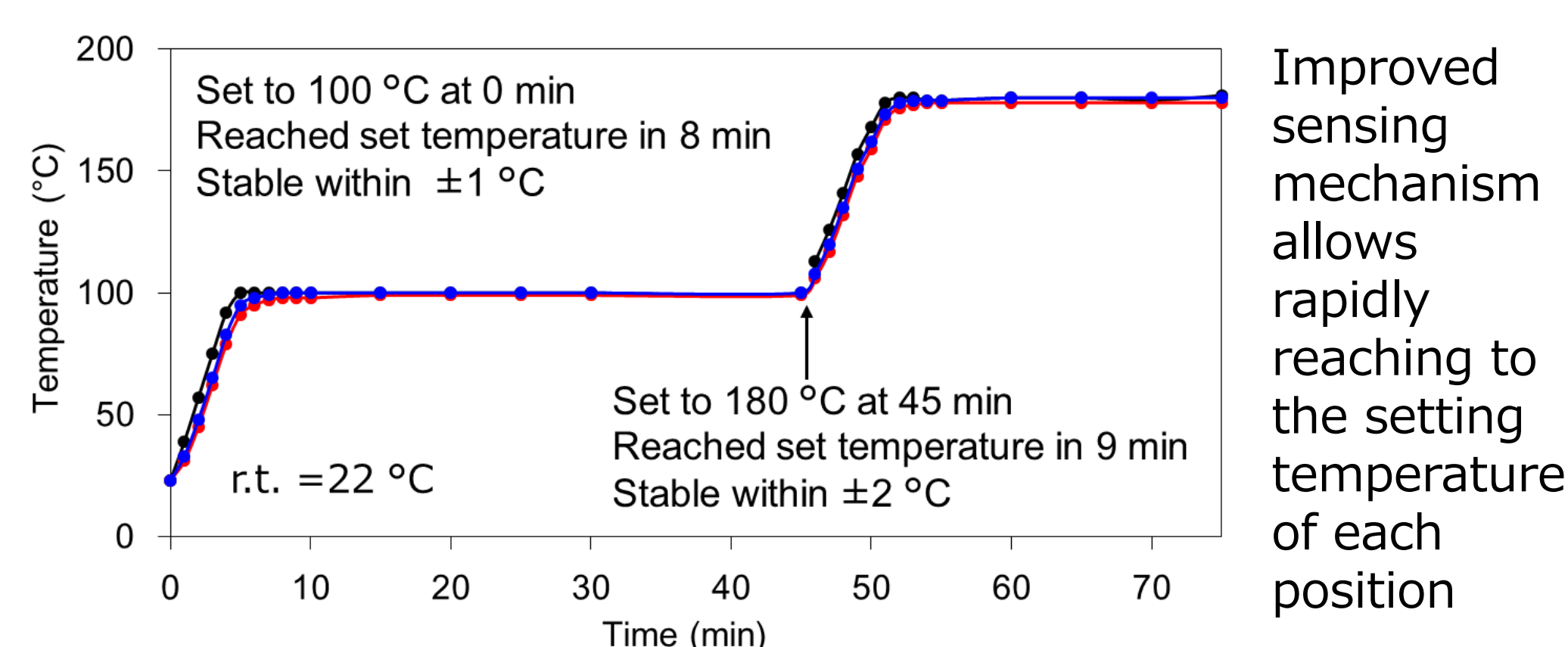


Summary

Recently demands on discovery and QC in drug development are increasing, hence facile and rapid hydrolyses of peptides and proteins are required. We have constructed a novel hydrolysis system which adapted to wide range of temperature without leakage of HCl fume. Two applications of AHST-16 are shown.

(1) A library consisting of cyclic octa-peptides as capturing molecules consisting of 24 natural and non-proteinogenic amino acids on the manner of "one peptide on one bead (OPOB)" focusing on drug discovery. The system are applied for rapid deconvolution of OPOB using MS/MS, of which methods has been improved based on the previously reported. (2) As an another application, super rapid amino acid analyses was performed, while 16 samples could be analyzed with rapid hydrolyses followed by the rapid determination by LC/MS using unique separation columns in a single day. Thus AHST-16 is useful for deconvolution immobilized peptides and for high throughput amino acid analyses in QC of peptides and proteins.

AHST-16 Hydrolysis system, reactions under sealing, higher efficiency & labor saving in acid hydrolysis:



Advantages

(1) Easy vial sealing **no-use by "open flame"** i.e. without burners; (2) For various chemical processes under sealed conditions. ; (3) Easy operation without skills; (4) Contamination free, semi-microscale, one touch sealing (just ca 1 min) after reaction excess reagents can be rapidly removed by nitrogen stream for ca 30 sec using the same apparatus.

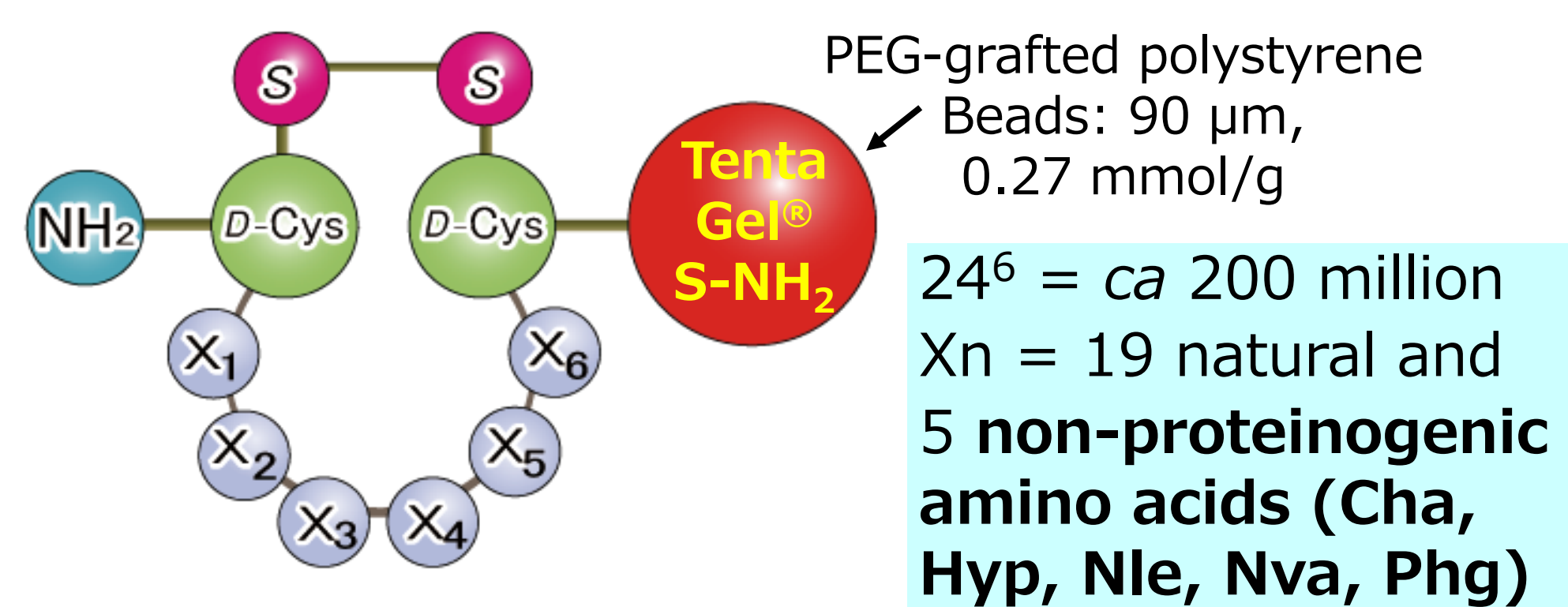
Part 1

Improved deconvolution method for cyclic peptides immobilized on Gel-type beads constructed for drug discovery tool

OPOB* has been prepared for drug discovery [*one single peptide immobilized on one bead, **LIBRARY of cyclic octa-peptides** (involving two **D**-Cys) immobilized on a gel-type resin (TentaGel®) which allows bioassays in an aqueous media

High quality cyclic peptide-beads as capturing molecules constructed with 24 amino acids and cyclized with two **D**-Cys = **Diversity 24⁶ = ca. 200 millions**; No inter-molecular disulfides; ca 100 µm particle; Peptide loading ca 80 pmole/bead; 2.3 million beads/g
Selected single bead are deconvoluted by partial hydrolysis with MS/MS

The present report is the improved method from previously reported: Nokihara et. al., *Amino Acids*, 48, 2492-2499, 2016. DOI 10.1007/s00726-016-2269-1

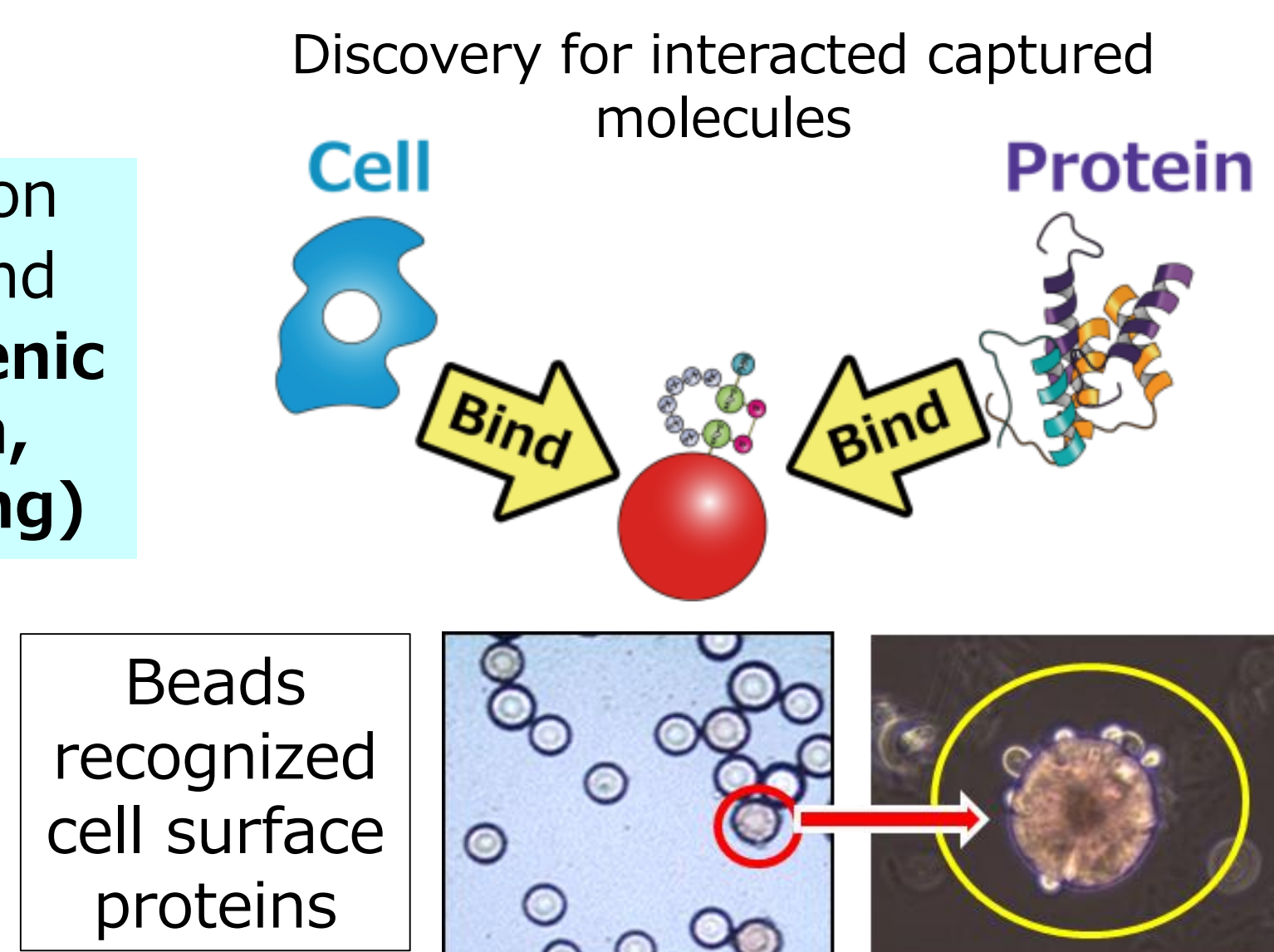


Construction of OPOB

Split & combine method, Deprotection and coupling are carried out using PetiSzyer® with LibraTubes®; On-resin cyclization in 10% DMSO (pH6.0) at room temperature for 4 days.
Detail: Nokihara et. al., *Amino Acids*, 48, 2492, 2016

OPOB: available from HiPep Labs (<http://hipep.jp/eng/?p=751>) as research products and/or Nakarai Tesque US.

OPOB is a useful tool for drug discovery

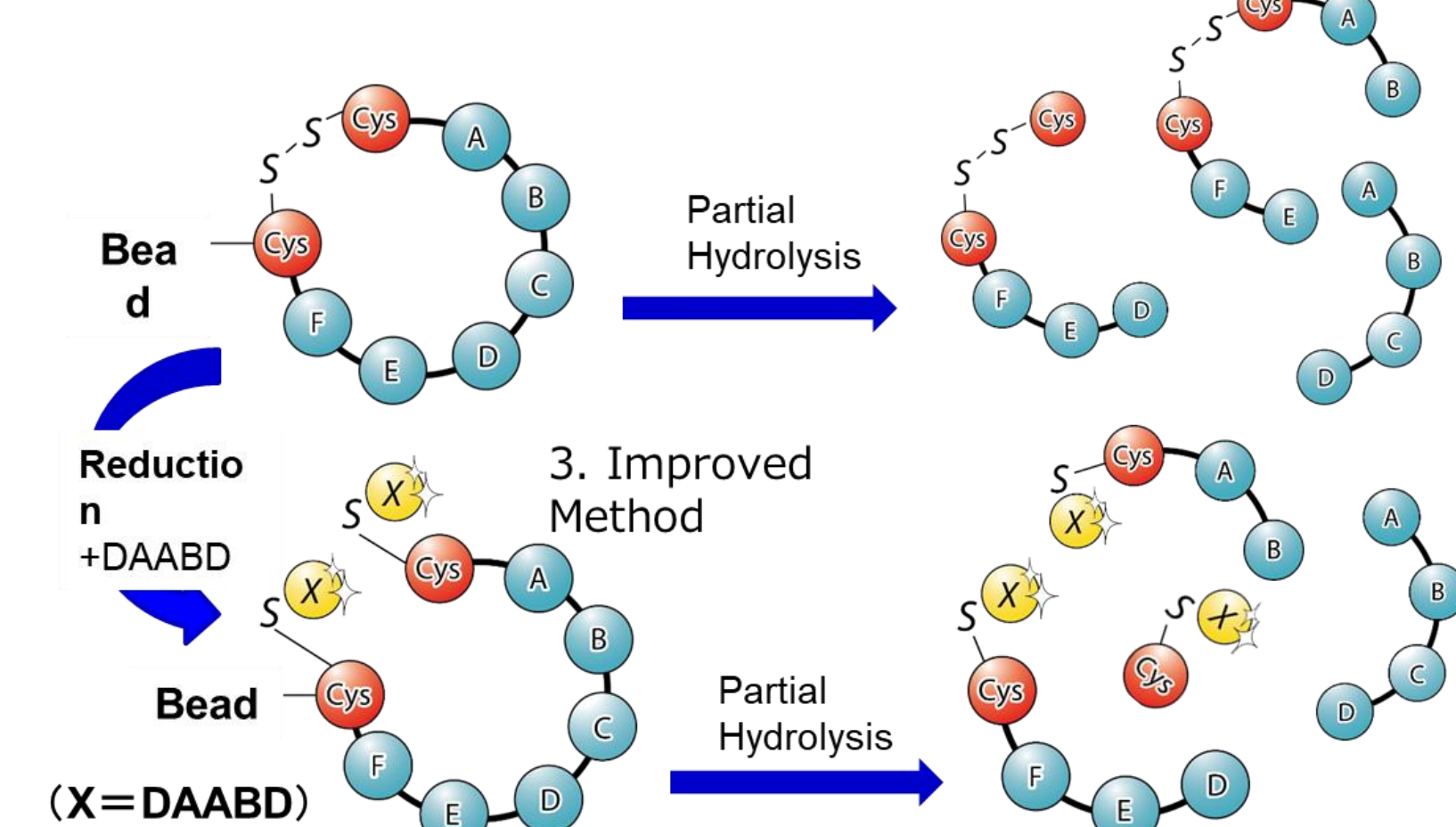


Screening → Deconvolution

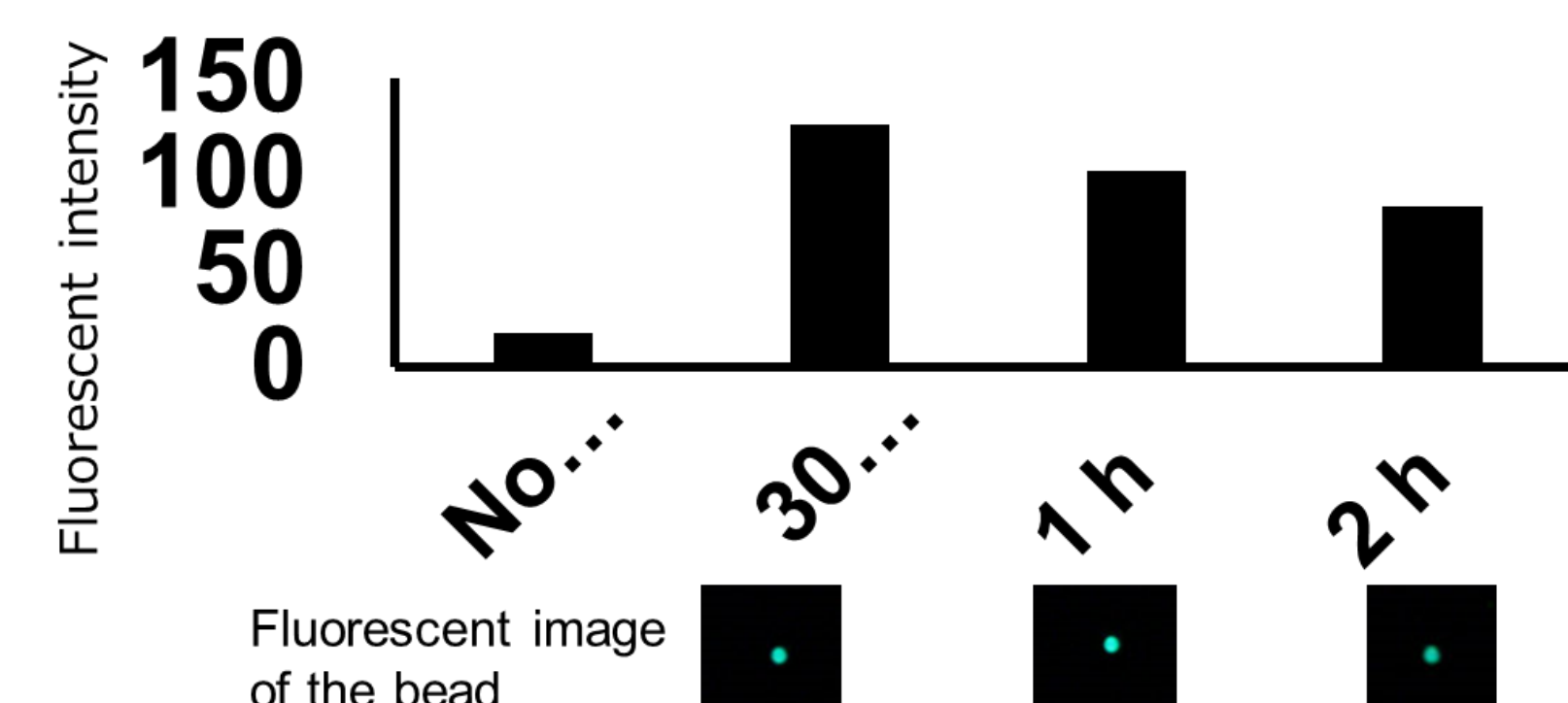
Partial hydrolysis with 5.7N HCl at 60 °C for 3 h by the use of the AHST-16 → LCMS/MALDI-TOF-MS/MS

Problems in deconvolution

1. LC-MS of partial hydrolysate of cyclotides on beads gave difficulties to identify the corresponding peaks
2. Termini were unclear since target was cyclic



Fluorescent intensity changes after hydrolysis of the labeled bead during hydrolysis (UV 375 nm)



Fluorescent intensity (UV375 nm) of the labeled bead was gradually decreasing depends on duration of hydrolysis = the immobilized peptide was gradually liberated (detached from a bead).

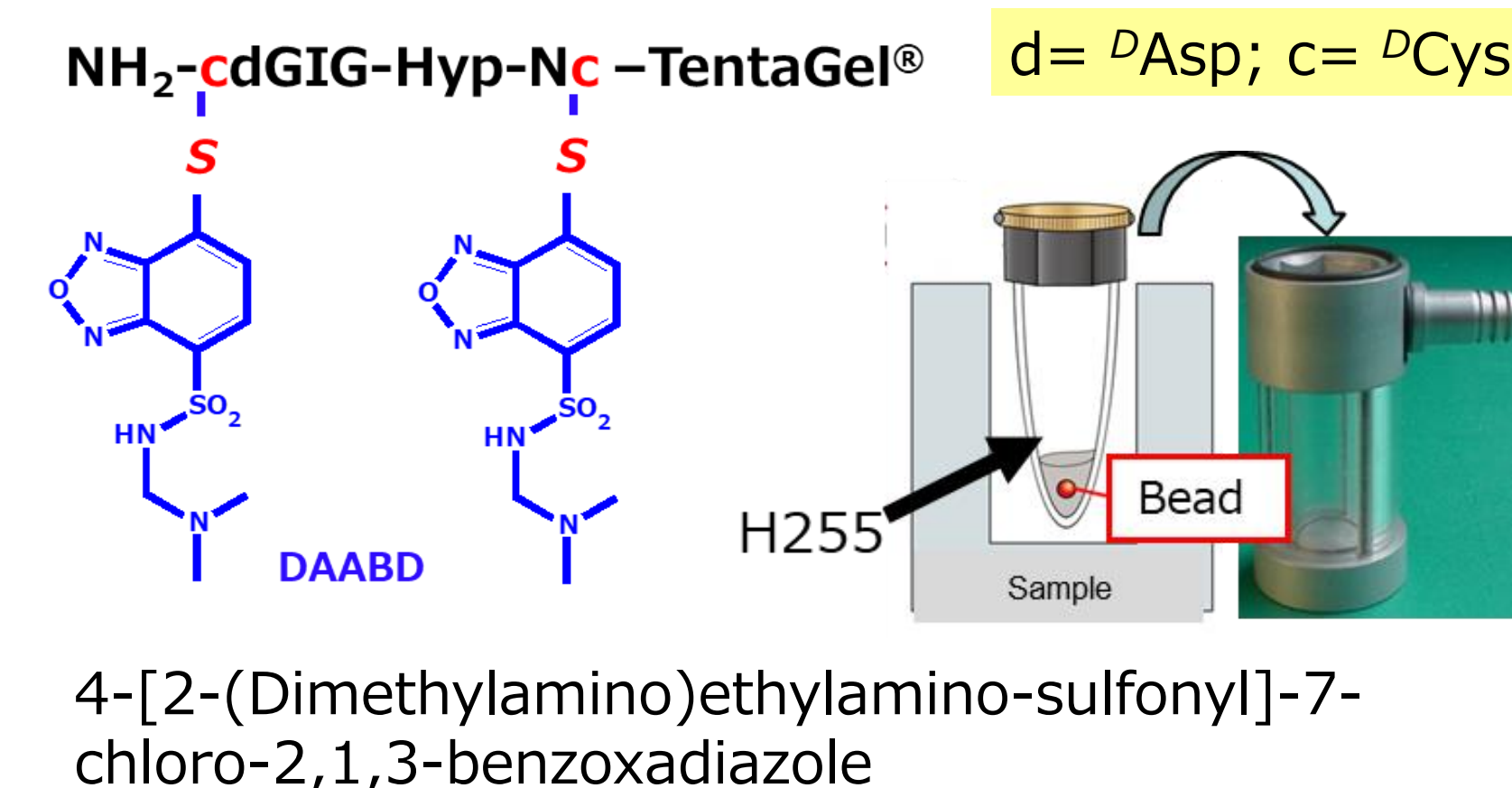
Hydrolysate: Analyzed by LC-MS [LC: Agilent 1100+ MS: HCTultra (IT-MS, Bruker)]

Conclusion of Part 1

By reducing and derivatizing for Cys-cyclotides with DAABD, the sensitivity on LC-MS was enhanced. Fluorescent labeling facilitated identification of the fragments generated by the partial hydrolysis, and sequence analyses were efficiently improved.

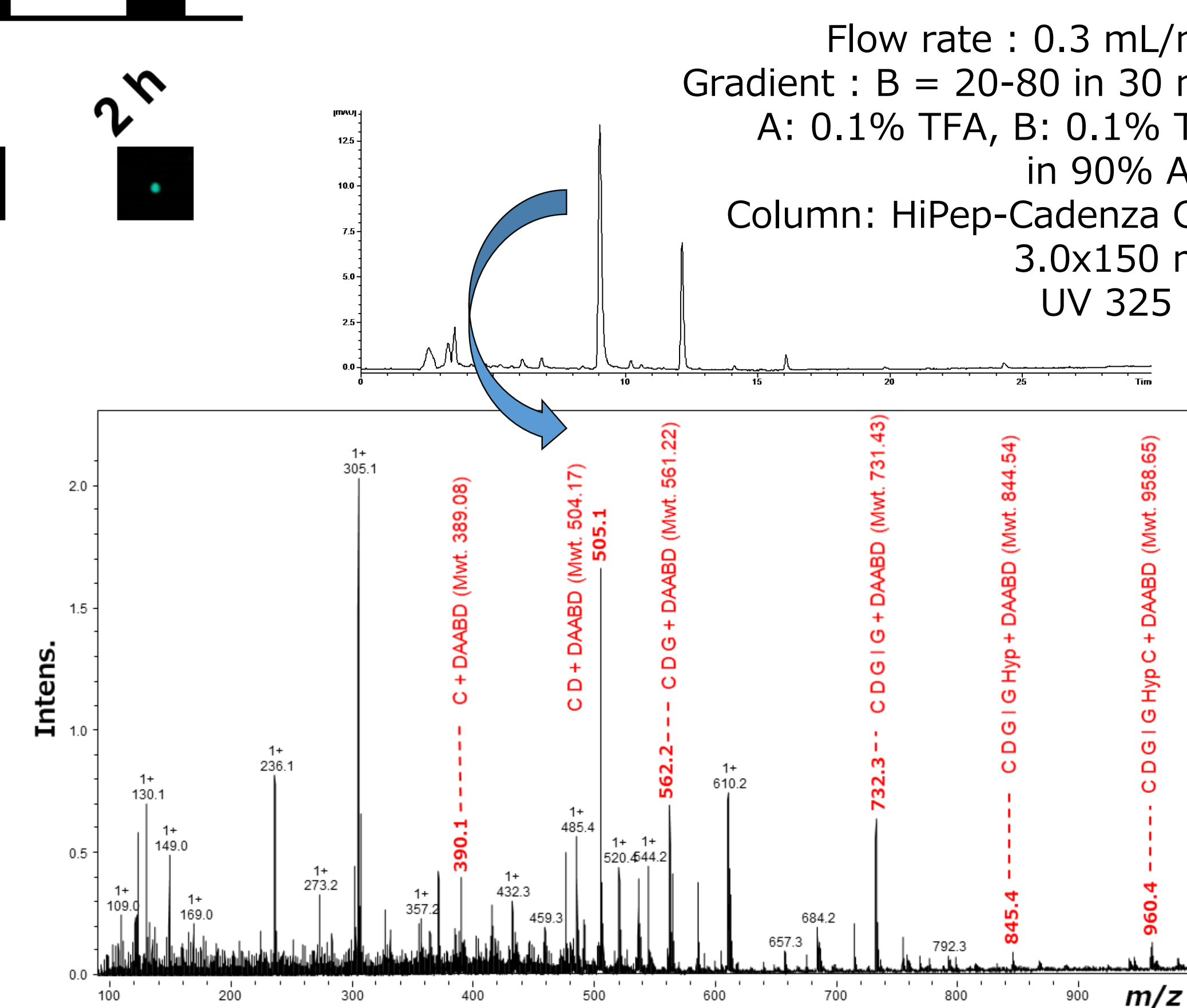
- ① Cyclic peptide beads were reductively derivatized with DAABD-Cl. The progress of the hydrolysis has been shown as fluorescence intensities of the bead was decreased. Thus the peptide on a bead was fragmented and released by the hydrolysis.
- ② The peptide has been partially decomposed by hydrolysis at lower temperature for shorter time.
- ③ In the case of a peptide consisting of Cys at both termini, fluorescent labeling of Cys residues resulted higher efficient deconvolution.
- ④ Hydrolysate of the OPOB-bead for 2hours several fragment peaks were observed at LC-MS analysis. The mass indicated in the spectrum gave agreement with the theoretical value of the corresponding fragment generated by hydrolysis.

Cysteine residues at both termini of cyclotide immobilized on bead were derivatized with DAABD*-Cl (fluorescent labelling agent) by the one-pot reaction (10 min)



Methods

1. A bead was reduced and labeled in 0.83 mM tris(2-carboxyethyl) phosphine), 0.14 M DAABD-Cl
2. Washed with MeOH, H₂O
3. Resulted bead was partially hydrolyzed in a vial H255 with 5.7 N HCl (20 µL) under sealing at 60 °C for 30 min, 1 h, 2 h using AHST-16

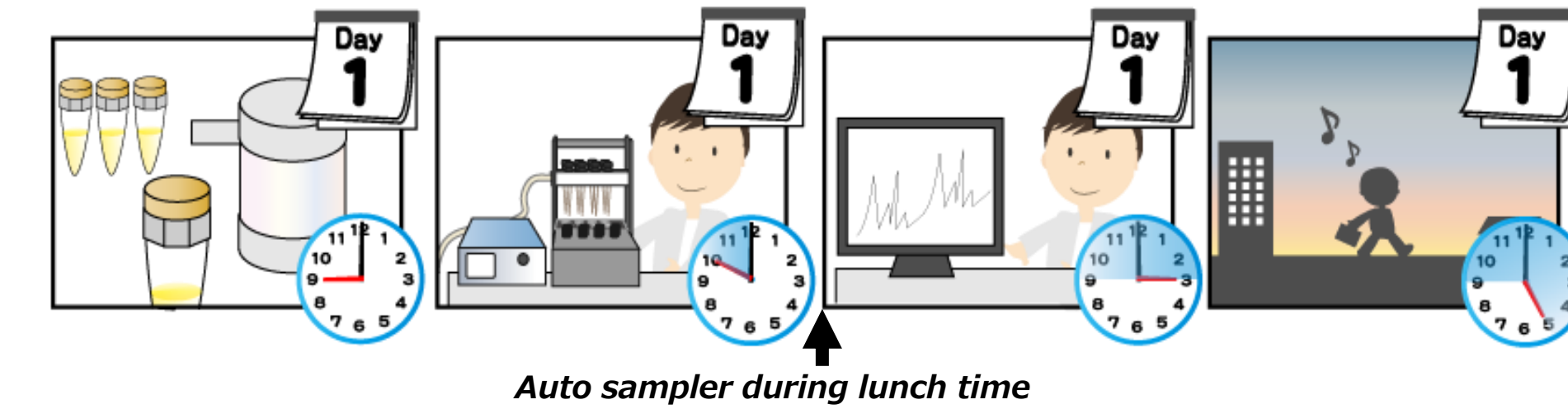


LC-MS of hydrolysate of the above DAABD-peptide: was labeled with DAABD (5.7 N HCl for 2 hrs)

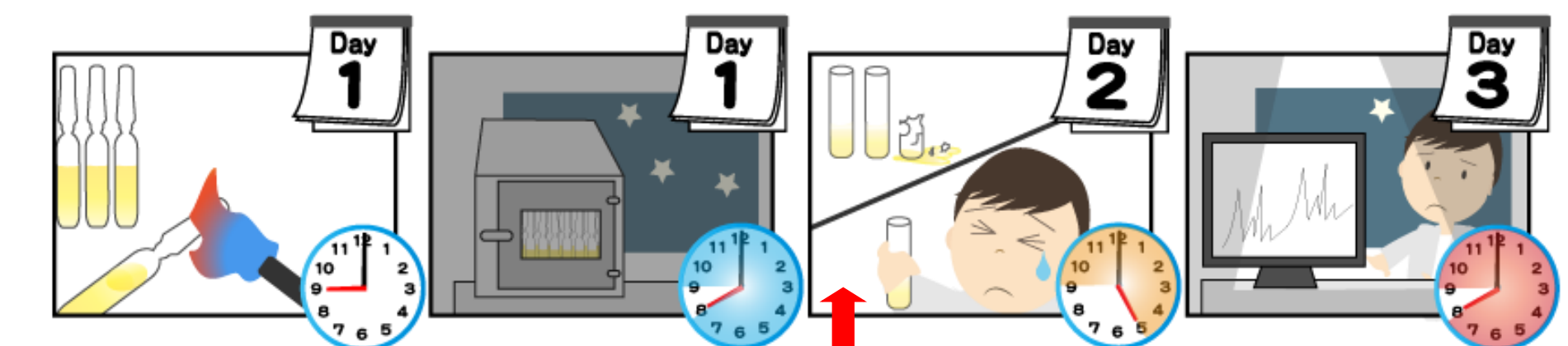
Part 2

High throughput Amino Acid Analysis by the Novel System without any derivatization
→ up to 16 Samples in a single day

Smart analyses Labor saving



Conventional and previous manner

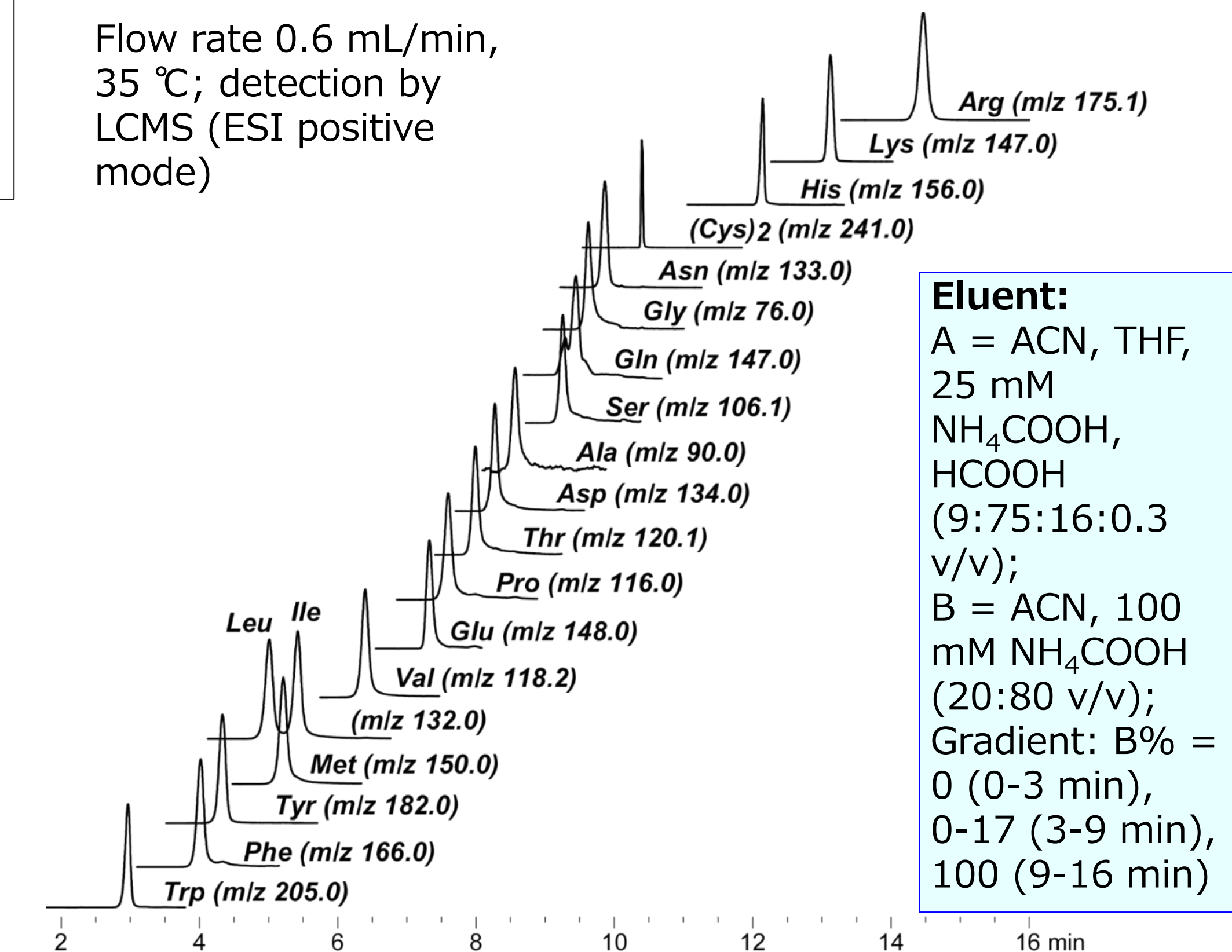


Vial broken, HCl fume destroys oven...Not completely hydrolyzed.....

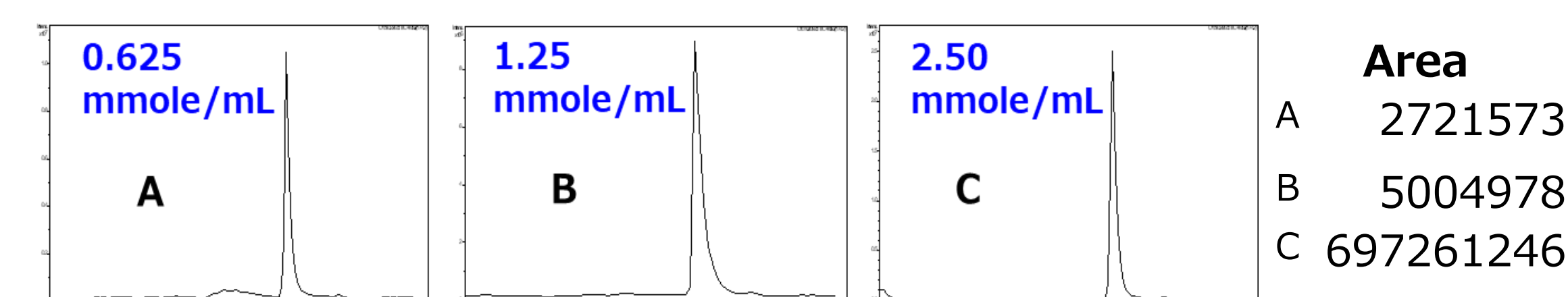
A typical separation profiles of standard amino acids without derivatization:

Column: Intrada Amino Acid 3 id x 100 mm

Flow rate 0.6 mL/min, 35 °C; detection by LCMS (ESI positive mode)



Quantitative analyses by a LC-MS



Conditions
Column: Intrada AminoAcid (150 x 3 id mm)
Flow Rate: 0.6 mL/min
Gradient :B%=20 (0-5 min), 35 (5-11 min), 100 (11-20 min)
A: 0.3% HCOOH in ACN, B: 100 mM Ammonium formate/ACN = 80/20, at 40°C; HCTplus, Bruker

AA	regression equation	R ²
F	y = 1E+09x+7E+08	0.9613
I	y = 9E+07x+2E+08	0.7582
L	y = 1E+08x+1E+08	0.9988
V	y = 1E+08x+7E+07	0.9554
E	y = 1E+08x+4E+07	0.9833
D	y = 5E+07x+1E+07	0.9770
N	y = 6E+07x+3E+07	0.9487
H	y = 2E+08x+5E+07	0.9979
T	y = 6E+07x+3E+07	0.9746
G	y = 1E+07x+3E+06	0.9817
Q	y = 2E+08x+2E+08	0.9509
C	y = 7E+07x+4E+07	0.9997
R	y = 4E+08x+3E+08	0.9097
S	y = 5E+07x+3E+07	1
P	y = 4E+08x+3E+06-8	0.9445
K	y = 2E+08x+1E+08	0.9411
M	y = 4E+08x+4E+08	0.9257

Conclusion of Part 2

Current difficulties in High Throughput analyses by MS → Determination: Gln vs Lys; Leu vs Ile; N- & C-termini : free or modified?; Gln & Lys give similar m/z, Leu Ile give the same m/z; High resolution MS (= costly instrument) + contaminants causes noise & disturbing interpretation

- ① High throughput amino acid analysis was performed by **AHST-16**, with the column **Intrada Amino Acid**, and **LC-MS**.
- ② Acid hydrolysis: **HCl:TFA = 2:1 v/v (166°C for 25 min) → Throughput at least 16 samples in a day**
- ③ The present AA-analysis gives useful information without costly instruments, such as **Amino acid analyzers**.

The KEYS for MS-de novo sequencing

- ① N-terminus: acetylation gave peak shift +42 Da
- ② C-terminus: methyl esterification gave peak shift +42 Da

