Investigation of Tryptic Protein Digestion in Microdroplets and in Bulk Solution

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Cite This: J. Am. Soc. Mass Spectrom. 2022, 33, 1238–1249

Abstract: Recent studies have shown that ultrafast enzymatic digestion of proteins can be achieved in microdroplet within 250 μs. Further investigation of peptides resulting from microdroplet digestion (MD) would be necessary to evaluate it as an alternative to the conventional bulk digestion for bottom-up and biotherapeutic protein characterization. Herein we examined and compared protein tryptic digestion in both MD and bulk solution. In the case of MD of β-lactoglobulin B, the preservation of long peptides was observed due to the short digestion time. In addition, MD is applicable to digest both high- and low-abundance proteins in mixture. In the case of digesting NIST 8671 mAb antibody containing a low level of commonly encountered host cell protein (HCP) PLBL2 (mAb:PLBL2 = 100:1 by weight), MD produced lower levels of digestion-induced chemical modifications of asparagine/glutamine deamidation, compared with overnight digestion. No significant difference between MD and bulk digestion was observed in terms of trypsin digestion specificity based on examination of semi- and unspecific-cleaved peptides. Our study suggests that MD, a fast digestion approach, could be adopted for bottom-up proteomics research and for peptide mapping of mAbs to characterize site-specific deamidation and glycosylation, for the purpose of development of biopharmaceuticals.

Keywords: enzymatic digestion, microdroplet reaction, antibody, deamidation

Introduction

Numerous studies in the past decade have shown that reaction rates can be dramatically accelerated if reactions are conducted in microdroplet rather than in bulk solution. The reaction acceleration has mainly been attributed to the special reaction environment at the air–liquid surface of microdroplet where partial solvation of the reagents, the presence of a strong electric field, enhanced concentration of solute, rapid desolvation, and surface excess charge accumulation could be the influencing factors to speed up the reaction. This “catalytic” feature allows microdroplet chemistry to have impactful applications including rapid chemical derivatization, reaction kinetic studies, high-throughput reaction screening, nanomaterial preparation, and the rapid, green and sustainable synthesis or degradation. However, most of these reactions involved small organic molecules, large molecule protein reactions in microdroplets are rare, except enzymatic digestion of proteins. The microdroplet digestion of proteins is very fast and completes in submillisecond time scale (e.g., within 250 μs).

Protein digestion is an important step for bottom-up proteomics research, as large proteins can be cleaved into small peptides that can be easily analyzed with mass spectrometry (MS). Traditional digestion of proteins is time-consuming and often needs overnight incubation of proteins and enzymes like trypsin. To accelerate enzymatic protein digestion, a variety of novel methods have been reported, including increasing the digestion temperature, adding organic solvents, applying microwave energy, using high intensity focused ultrasound or employing microchip reactor. Microdroplet digestion appears to be an ultrafast approach in this regard. However, detailed investigation of microdroplet protein digestion has not been done, including the examination of peptide miscleavage, cleavage specificity, and chemical modification involved in digestion.

Monoclonal antibodies (mAbs) and related products are the fastest growing class of human therapeutics in the pharmaceutical industry, and many mAb products have been approved for treatment of different pathologies including cancers, and immunological disorders. Most therapeutic mAbs are a class of recombinant proteins that are susceptible to a variety of enzymatic or chemical modifications, also called post-translational modifications (PTMs), during cell culture, purification, formulation and long-term storage. Common PTMs previously identified include glycosylation, deamidation,
Scheme 1. Workflows of Microdroplet and Bulk Digestions in This Work

Isomerization, and oxidation, incomplete disulfide bond formation, glycation, N-terminal glutamine cyclization, and C-terminal lysine processing. Deamidation is one of the most common chemical modifications observed during the degradation processes of mAbs, which may result in changes in physiochemical properties, such as hydrophobicity, charge, and secondary or tertiary structure, and may lower the thermodynamic or kinetic barrier to unfold. This may affect the stability, binding affinity, half-life, and efficacy of mAb products, potentially causing immunogenicity. Therefore, characterization and control of such modifications is important for the quality control of antibody drugs throughout the product life cycle. Currently, peptide mapping, a bottom-up strategy, via reversed-phase liquid chromatography coupled with mass spectrometry (LC-MS) is considered as a gold standard for monitoring and characterization of site-specific modifications including deamidation that may arise during production, processing, or storage. However, to generate a peptide map, the therapeutic protein must first be digested into its constituent peptides through enzymatic reaction in which the sample needs to be exposed to various buffers, even elevated temperatures for up to 24 h to reach the maximum digestion efficiency. Unfortunately, the conditions that are amenable for effective digestion are often similar to conditions that promote unwanted modifications. For example, it has been reported that asparagine deamidation is promoted by conditions such as elevated temperatures or high pH and are further exacerbated by exposure to these extreme conditions for extended incubation time. Thus, it is often challenging to find a balance between maximum digestion efficiency and minimizing artificial modifications. Considering the ultrafast speed of microdroplet protein digestion, we reason that chemical modifications on antibody peptides could be reduced if microdroplet digestion is used to replace traditional digestion protocols.

In this study, we evaluated the performance of microdroplet protein digestion including digestion efficiency, sequence coverage, peptide cleavage specificity and deamidation resulted from digestion. Both model protein β-lactoglobulin B and antibody mAb were tested. In general, in comparison to bulk solution digestion, long peptides tend to be well preserved in microdroplet digest probably due to the much shorter digestion time and room temperature used. Equally importantly, deamidation is reduced. Our results suggest that microdroplet-assisted enzymatic reactions could serve as an alternative to traditional digestion method commonly used in bottom-up proteomic studies, as well as for the biotherapeutic protein characterization.

EXPERIMENTAL SECTION

Chemicals. Monoclonal antibody reference material 8671 (NIST 8671 mAb) was obtained from the National Institute of Standards and Technology (NIST, Gaithersburg, MD). Phospholipase B-like 2 protein (PLBL2, CHO-S-purified) was purchased from Immunology Consultants Laboratory (Portland, OR). β-Lactoglobulin B from bovine milk, trypsin (sequencing grade), urea (electrophoresis grade), ammonium bicarbonate (biotra grade), dithiothreitol (DTT, biotra grade), and iodoacetamide (IAM, biotra grade) were all sourced from Sigma-Aldrich (St. Louis, MO). Peptide gp 100
For bulk digestion, an equal amount of protein and enzyme was mixed (25 μL of 1 μg/μL β-lactoglobulin B and 25 μL of 0.05 μg/μL trypsin) and incubated at 37 °C overnight (14 h). FA was added to 1% (v/v) to quench the digestion. The collected digests from both microdroplet and bulk digestions were vacuum-dried and reconstituted with 60 μL of H₂O/ACN/FA (50:50:1, v/v). Fifteen microliters of peptide TKPR (355.9 μM) was added as the internal standard for nanoESI-MS analysis. The injection flow rate was 2 μL/min, and +3.5 kV was applied for nanoESI ionization. The temperature of the MS inlet was 250 °C. A Thermo Scientific Q Exactive mass spectrometer (San Jose, CA) was used with resolution set to 140,000 for the MS analysis.

**Microdroplet Digestion of a Mixture of mAb and PLBL2.** One milligram of NIST 8671 mAb was spiked with 10 μg of PLBL2 to mimic an antibody sample containing an impurity protein. The sample was then, diluted to 1 mg/mL mAb and 0.01 mg/mL PLBL2 with 50 mM ABC buffer (pH 8.0) containing 8 M urea. The denatured protein mixture was reduced with 20 mM DTT at 37 °C for 1 h and alkylated with 40 mM IAM at room temperature in the dark for 30 min. A second aliquot of DTT (20 mM) was added again to quench extra amount of IAM. The alkylated protein sample was loaded into Amicon Ultra-0.5, 3 kDa filters (Millipore Sigma), then centrifuged at 14,000 g for buffer exchange to remove small molecule reagents like urea, DTT and IAM. The concentrated protein solution was diluted with 100 mM ABC buffer to 1 μg/μL digestion. As shown in Scheme 1, for microdroplet digestion of β-lactoglobulin B, 200 μL of 1 μg/μL protein in ABC buffer was added into one syringe, and 200 μL of 0.05 μg/μL trypsin in 0.1% acetic acid was loaded into the other syringe. The injection flow rate of both syringes was 5 μL/min. The protein and enzyme were mixed via a T-mixer and then delivered to a homemade sonic spray ionization (SSI) emitter for generating microdroplets of the reaction mixture. The connection capillary between T-mixer and emitter was 8 cm long with 98 μm ID (the dead volume = 0.6 μL). A 1.5 mL Eppendorf tube vial containing a quenching solvent of 50 μL of H₂O/ACN/FA (50:50:1, v/v) was used to collect the sprayed microdroplets (the distance between the emitter tip and the quenching solution in the vial was about 2 cm). The SSI emitter was composed of an inner fused-silica capillary (98 μm ID, 200 μm OD) and a coaxial outer fused-silica capillary (250 μm ID and 355 μm OD) and N₂ gas of 120 psi was used as the spray sheath gas to assist formation of microdroplets (no high voltage was applied). The coaxial capillary outlets were flush to reach the optimum spray geometry with smallest droplet sizes previously measured. On the basis of the previous report of microdroplet digestion employing an emitter with the same capillary dimensions, the same nebulizing gas pressure and the same spray distance as used in this work, the microdroplet digestion time of this experiment (i.e., the flight time of the sprayed microdroplets between the sprayer and the quenching solution) was estimated to be 250 μs (the same as previously reported). The sprayed microdroplet was collected for 5 min for subsequent nanoESI-MS analysis.

During the microdroplet digestion experiment, as a control experiment to assess the contribution of bulk digestion inside the short connection capillary after mixing of protein and trypsin via the Tee, the outlet end of the connection capillary was directly immersed in the quenching buffer of H₂O/ACN/FA (50:50:1, v/v) contained in a vial (setup shown in Scheme S1). Fifty microliters of 1 μg/μL protein in ABC buffer was loaded into one syringe. Then, 50 μL of 0.05 μg/μL trypsin in 0.1% acetic acid was loaded into the other syringe. The flow rate of both syringes was 5 μL/min. The protein and enzyme were mixed via a Tee and collected for 5 min for nanoESI-MS analysis.
from the first event. Ions were generated using a sheath gas flow rate of 35, an auxiliary gas flow rate of 10, a spray voltage of 4.0 kV, a capillary temperature of 300 °C, an auxiliary gas temperature of 100 °C and an S-Lens RF level of 50. Resolution was set at 70 000 (AGC target of 3e6) and 17 500 (AGC target of 5E4) for survey scans and MS/MS events, respectively. The maximum ion injection time was 100 ms for survey scan, and 100 ms for the other scans. The dynamic exclusion duration of 8 s was used with a single repeat count.

**Data Analysis.** Raw files were searched using MaxQuant software (version 2.0.3.1) equipped with the Andromeda search engine (Max Planck Institute, Martinsried, Germany). The first search peptide mass tolerance was set to 20 ppm, and the main search peptide mass tolerance was 4.5 ppm. Trypsin was set as the digestion enzyme with a maximum of two miscleavages. Carbamidomethylation was set as a fixed modification, while oxidation (on methionine M), acetylation (protein N-term), and deamidation (occurring to either asparagine N or glutamine Q) were set as variable modifications. Spectra were searched against the sequence of NIST 8671 mAb light chain and heavy chain plus the sequence of PLBL2. The “match between runs” feature was used with a matching time window of 0.7 min and an alignment time window of 20 min. Target decoy analysis was performed by searching a reverse database with a PSM FDR of 1% and protein FDR of 5%. Label-free quantification (LFQ) was performed using the MaxLFQ algorithm in MaxQuant software (MaxPlanck) according to default LFQ parameters with minimum ratio count set as 2, normalization type as classic, minimum number of neighbors equal to 3 and average
number of neighbors equal to 6. A separate database search was performed with digest mode set to semispecific and unspecific, respectively. The MaxQuant output tables were used for further analysis and visualization of data.

## RESULTS AND DISCUSSION

To evaluate the performance of microdroplet protein digestion, we started with trypsin digestion of a single protein first and β-lactoglobulin B was chosen as the test sample. We used an online mixing setup as shown in Scheme 1 for microdroplet digestion, for the reason that the online mixing would minimize the contribution of protein digestion in bulk phase before performing microdroplet digestion. As shown in the microdroplet digestion setup (Scheme 1), the residence time of protein and trypsin in the connection capillary before spray was estimated to be 4 s, based on the capillary dead volume of 0.65 μL and the mixed solution flow rate of 10 μL/min. In the control experiment (Scheme S1 and Figure S1), online mixing of β-lactoglobulin B and trypsin for 5 s led to little digestion of the protein and the acquired MS spectrum (Figure S1b) showed intact protein ions (+10 to +21 ions) along with two peptide ions at m/z 419 and m/z 858 were detected. m/z 419 and 858 corresponded to +2 ions of ALPMHHR and LSFNPTQLIEQCHI, respectively, two peptides from the C-terminal of β-lactoglobulin B, which were also observed in the MS spectrum of intact protein (Figure S1a).

Using the Scheme 1 setup for microdroplet digestion of β-lactoglobulin B, we used two syringe pumps to deliver protein and enzyme solution separately at 5 μL/min flow rate. The reactants were mixed online via a T-tee before spray. Microdroplet digestion was quenched immediately by the quenching solution after reaching the collection vial. Overnight bulk digestion (at 37 °C) of β-lactoglobulin B was conducted for comparison. After digestion, both digestes were vacuum-dried and reconstituted in solvent with peptide TKPR being spiked as an internal standard (IS) for nanoESI-MS analysis. As shown in Figure 1a, after digestion, numerous peptide fragments were observed while signals corresponding to the intact protein could no longer be seen in both microdroplet and bulk digestion, indicating the tryptic digestion in both conditions was complete. Moreover, by manually searching fully cleaved peptides (Figure 1a inset) based on the sequence of β-lactoglobulin B, the same number of fully cleaved peptides were identified both in microdroplet and bulk digestion, leading to the same sequence coverage (Figure 1b).

Interestingly, there were some peptides (highlighted in red, Figure 1a inset) showing higher abundance in microdroplet than bulk digestion. For example, top two long peptide VAGTWWYSLAMASDSSLDAQSPAPLRL and YLLFCMENS-AEPEQSLACQCLVR were detected with much lower intensities in overnight bulk digest (2.64 × 10^7 and 5.63 × 10^6) than in microdroplet digest (5.18 × 10^8 and 2.51 × 10^8) (identified peptide intensities are summarized in Table S1). This result indicates that long peptides might suffer hydrolysis in solution with long incubation time and elevated temperature at 37 °C, while they could be well preserved during short microdroplet digestion period. This finding is consistent with the previously reported proteome studies in which long digestion time resulted in reduced number of identified peptides and low protein sequence coverage.35–37

In addition, we also examined the digestion-induced chemical modification of asparagine deamidation of our detected peptides (Figure 1a) in microdroplet and bulk digestes. Deamidation as an important PTM during protein degradation, has been widely studied and known to be accelerated at high temperatures and high pH values.38 Indeed, one peptide from β-lactoglobulin B, WENGECQAQK, had much higher deamidation level observed in the bulk overnight digest. As shown in Figure 1c, only native form of WENGECQAQK was detected at m/z 1121 from MD (theoretical mass m/z 1121.46803, mass error: 1.1 ppm); in contrast, the deamidated form, WEDGECQAQK with 1 Da shift (m/z 1122.45008, theoretical mass m/z 1122.45205, mass error = 1.8 ppm), was observed in bulk digest (Figure 1d). This result indicated that the ultrafast microdroplet digestion (∼250 μs) at room temperature also tends to prevent the digested peptides from deamidation, besides preserving long peptides.

Because reproducibility is important for protein digestion, in this study, we repeated the microdroplet digestion with three individual runs using the online mixing setup described above (Scheme 1). The nanoESI MS spectra for triplicate microdroplet digestions (a–c) and triplicate bulk overnight digestions (d–f) are shown in Figure S2. Similar profile across digestion replicates were observed for each digestion condition. In general, microdroplet digestion showed high reproducibility between three individual runs in terms of the sequence coverage. For most of the fully cleaved peptides, the standard deviation among three digestion replicates is acceptable (Figure S3).

In our microdroplet digestion setup (Scheme 1), to reduce the loss of peptides from collection of the sprayed microdroplets, the collection vial was covered with a lid and the emitter tip was inserted into the vial through the lid via a predrilled hole. As a test, standard peptide KVPRNQDWL solution was sprayed and collected using the setup shown in Scheme 1. By comparing the sprayed peptide signals before and after the spray, we measured the collection recovery to be 105% (details shown in Table S2), indicating that there was no notable peptide loss during microdroplet spray collection process.

After examining microdroplet digestion with β-lactoglobulin B as an example, we moved forward to investigate the microdroplet digestion of proteins in mixture. No digestion of multiple proteins in microdroplet was examined before. In our experiment, NIST 8671 mAb was mixed with HCP protein PLBL2 at 100:1 ratio to mimic a protein mixture where one protein is dominant, and the other one is a minor species. PLBL2 is a 66KDa catalytic HCP protein that copurifies frequently in mAb preparations produced in Chinese hamster ovary (CHO) cells, which might cause the unwanted immunogenic effects in patients and affect the drug stability due to its enzymatic degradation of polysorbates 20 and 80, common adjuvants present in formulated drug substance (DS).39,40 Therefore, PLBL2 needs to be removed from mAb preparations down to acceptable levels, and therefore, it is necessary to monitor the presence of PLBL2 in IgG antibodies. In this study, we attempted to examine how the microdroplet digestion would occur to a low-level PLBL2 protein and a high-level concentration of large mAb antibody. On the other hand, we evaluated if chemical modifications to antibody like asparagine/glutamine deamidation could be reduced during the short digestion time of MD.

Microdroplet digestion and bulk digestion of the mixture of NIST 8671 mAb and PLBL2 were conducted as described in Experimental Section. Both microdroplet digest and bulk
overnight digest were prepared as 3 digestion replicates and then subject to LC/MS analysis. Following LC/MS/MS analysis, peptide identification and quantification were performed to evaluate the quality of the digested samples between the two different digestion conditions. The digestion efficiency was studied in terms of the determination of sequence coverage, missed cleavage, semispecific cleavage, and nonspecific cleavage as the metrics. As shown in Figure 2a, comparable sequence coverage of antibody heavy chain was attained for both microdroplet and bulk digestions. A slightly higher sequence coverage of antibody light chain and lower abundance protein PLBL2 in microdroplet was observed compared with bulk digestion. Upon further investigating the identified peptides containing zero, one, or two missed trypptic cleavages (Figure 2b–d), microdroplet and bulk overnight digestion showed comparable trypsin digestion efficiency for mAb heavy chain. However, for mAb light chain and low abundance PLBL2, microdroplet showed less miss cleaved peptides than bulk.

Aside from the missed cleavages, we also checked the semispecific cleavage (a specific cleavage at one terminus and a nontryptic cleavage site at the other end) and nonspecific cleavage (enzymatic cleavage of C-terminal to a residue other than Lys or Arg for both termini), since microdroplet digestion was conducted in a new reaction environment with unknown reaction mechanism which may affect the enzyme specificity. As shown in Figure 2e, bulk overnight digestion showed higher level of semispecific cleavages with 156 peptides identified, whereas the number of identified semispecific peptides from microdroplet digest was 115. This trend continued when comparing nonspecific cleavage in Figure 2e, a lower-level nonspecific cleavage was observed in microdroplet digestion with 39 peptides identified, whereas 74 nonspecific cleaved peptides were found from bulk overnight digestion. All these results suggested that microdroplet digestion has comparable digestion efficiency with bulk and no significant increment of semispecific cleavage, as well as nonspecific cleavage, was observed.

It is noteworthy that a significant number of semispecific and nonspecific trypptic peptides were identified in the case of antibody sample digestion. However, most of these peptides were of relatively low-abundance compared to fully trypptic peptides. For example, as shown in Figure S4a, four semitryptic peptides were detected corresponding to fully trypptic peptides VYACEVTHQGLSSPVTK (amino acid 190–206) from light chain. However, the relative abundance of those semitryptides were lower than 1%. Similar result was observed for another peptide TPEVTCVVVDVSHDEPEVK (amino acid 259–277) from heavy chain (Figure S4b). Although increased number of semitryptic peptides were detected, the overall relative abundance corresponding to the fully trypptic peptides were low. The intensity distributions of semispecific and unspecific peptides are shown in Figure S4c and d, whereby the major peptides were detected at low abundance (lower than 1 × 10^4). Indeed, in previous studies, nonspecific trypsin cleavages have been frequently reported and showed a high percentage especially in standard protein digest.51,62 Several potential sources for generating partially trypptic peptides has been reported, including possible chymotrypsin contamination in trypsin, pseudotrypsin activity due to partial autolysis of
trypsin, proteolytic products due to endogenous proteases, and in-source fragmentation of fully tryptic peptides.\(^{62}\) Besides, it has been shown to be related to a number of digestion conditions and predigestion treatment.\(^{63}-^{66}\) In this work, to prevent the autolysis of trypsin before spray, trypsin was reconstituted in 0.1% acetic acid. Although mildly acidic environment was often recommended by the manufacturer to prevent trypsin autolysis, a significantly increased level of nonspecific cleavages during the trypsin digestion process has been reported before.\(^{67}\)

In addition, we quantitatively compared each fully cleaved peptide based on the peak intensities reported from Maxquant. Interestingly, the abundance of different fully cleaved tryptic peptides was found to be quite distinct under the two digestion conditions. For example, as shown in Figure 3a and b, several long peptides showed much lower intensities in bulk overnight digest than in microdroplet digest, such as VTNMDPADTATYYCAR, WQQGNVF51SVMHEALHNHYTQK, ESGPALKPTQTLTETCSTFSFSLTAGMSVWIR, TPEVTCVVDVSHKPEVK, and TTHCPCPAPELG-GPSVLFPPPKPK. It is likely that these peptides are long and more susceptible to hydrolysis during bulk overnight digestion at 37 °C, whereas microdroplet digestion could significantly minimize the hydrolysis because of the fast digestion speed at room temperature. However, the other long peptides from the mAb heavy chain, such as DMIFNFY-
Figure 4. Averaged intensities of fully cleaved peptides identified from PLBL2 under different digestion conditions. Standard deviations from the reported mean shown are for $n = 3$ digestion replicates.

Figure 5. Averaged relative abundances of (a) deamidation for mAb and PLBL2 and (b) glycosylation for mAb heavy chain N300 under different digestion conditions. Averaged relative abundance was calculated as the intensity ratio of one modified peptide to the intensity sum of modified and native peptides. Peak intensities of native and modified peptides were recorded after manually searching of $+2$ charge forms of each peptide from MS1 spectra with 5 ppm mass accuracy tolerance. Standard deviations from the reported mean shown are for $n = 3$ digestion replicates.

FDVWGQGTTVTVSSASTK and GFYPSDIAVWESNGQPENNYK, showed higher abundance in bulk overnight digest than microdroplet.

The same trend was observed for mAb light chain (Figure 3c) wherein long peptides like SGTASVCLLLNFFPR, VYACEVTQHQLSLSPVTK, VDNALQSGNSQESVTEQDSK, and FSGSGSTETTLTISSLQPDFAFYYYCFQGSGYPTFGGGTK, were detected with lower intensities in bulk overnight digest than microdroplet digest. For low abundance protein PLBL2 (Figure 4), most of the fully cleaved peptides showed a relatively higher intensity in microdroplet than bulk, indicating that microdroplet might have a slightly higher digestion efficiency of low abundance proteins. It has been reported that the digestion bias exists between low and high
abundance proteins due to the fact that the rate of trypsin digestion is dependent on the concentration of each individual protein in a mixture. Common strategies to address the abundance challenge include affinity depletion and enrichment of proteins with antibody arrays or ligand libraries and prefractionation of proteins. Our microdroplet and bulk digestion of PLB2, the sequence coverage was about 40% (Figure 2a). To further improve the sequence coverage of spiked PLB2, the prefractionation or affinity enrichment could be the solution.

Upon further investigation of peptides that contain asparagine deamidation site as shown in Figure 6a. For example, deamidation sites, such as N38S and N469 from PLB2 and N136 and N151 from the light chain, as well as N300, N289 (279), N387, N364, N78Q79, and N318 from the heavy chain, were identified and relative levels are quantified, as shown in Figure S5a (denotation of N289/279 means either N289 or N279 underwent deamidation, the same for N78Q79). Overall, much higher level of deamidation was observed in bulk overnight digest than microdroplet. As the rate of deamidation also depends on the adjacent amino acid residue's size (smaller size reduces steric hindrance for the initial cyclization reaction of the deamidation process), the level of deamidation varies among different peptides. For instance, Gly is considered as the most favorable amino acid promoting deamidation, followed by Ser, Thr, and Asn, which is in line with the detected deamidation rate displayed in Figure S5a in which high deamidation level site like N385 from PLB2, N387, and N318 from heavy chain are followed by Gly, whereas N136 from light chain is adjacent to Asn. All these results indicated that reducing the digestion time without reducing digestion efficiency is preferable to minimize the digestion-induced deamidation.

Interestingly, EEOY100NSTYR containing one glycan site at N300 was detected with slightly higher deamidation rate in bulk overnight than microdroplet (Figure 6a). Upon further investigation of glycans, deamidated and unmodified peptides of EEOY100NSTYR in Figure 6b, we found that the relative abundance of unmodified peptides in microdroplet and bulk were similar, indicating deamidation rate is relevant to the glycan loss. Indeed, this result is consistent with the previous report that glycan loss in solution is accompanied by the conversion of Asn to Asp. Thus, a lower level of deamidation rate observed in microdroplet digestion might prevent glycan shedding that occurs during overnight digestion.

Finally, glycosylation was also assessed for the glycan profiles on heavy chain N300 site, as shown in Figure 6b. Most abundant glycan forms including complex biantennary oligosaccharides containing 0 to 2 nonreducing galactoses with fucose attached to the reducing end of N-acetylgalcosamine (A2G0F, A2G1F, A2G2F, A1G0F, A1G1F), along with high mannose structures (M5), were searched. The ratio of the relative abundances of most glycan forms was consistent between bulk and microdroplet digests, suggesting that microdroplet digestion could be applied to monitor PTMs for mAb.

In addition, in the microdroplet digestion experiments described above, the flow rate for protein injection was low. We further explored the possibility of using a high flow injection rate like 100 μL/min for each pump. In our test, 100 μL of 1 μg/μL β-lactoglobulin B in ABC buffer was loaded in one syringe and 100 μL of 0.05 μg/μL trypsin in 0.1% acetic acid was loaded in the other syringe for microdroplet spray (using the apparatus shown in Scheme 1). The flow rate of both syringes was 100 μL/min and N2 was set at 120 psi. The microdroplet was collected into a vial containing a quenching solution of 50 μL of H2O/ACN/FA (50:50:1, v/v) for just 10 s MS spectrum (Figure S5) showed β-lactoglobulin B was still well digested under such a high flow rate. The tolerance of high sample flow rate for microdroplet digestion would further speed up the total collection procedure.

Overall, the data presented in this study demonstrate that microdroplet digestion has the potential to significantly speed up the bottom-up LC/MS analysis of mAb, stemming from the efficient digestion, even for the low abundant proteins. Moreover, ultrafast digestion performed at room temperature allows to minimize artificial modification like asparagine/glutamine deamidation.

CONCLUSIONS

Fast enzymatic reactions of proteins and antibodies in microdroplet were investigated with detailed peptide analysis in this study. Some interesting features of MD were found. First, MD is fast, resulting in the preservation of long peptides in digest. Second, fast microdroplet digest could reduce deamidation of asparagine or glutamine residues, in comparison to bulk digestion. Third, MD is applicable to digest both high- and low-abundance proteins in a mixture. Fourth, no significant difference between MD and bulk digestion was observed for trypsin digestion specificity based on examination of semi- and unspecific-cleaved peptides. Fifth, similar glycan profiles on heavy chain N300 site were observed in both microdroplet and bulk digestions. A lower level of deamidation degradation was found in microdroplet than in bulk. Sixth, microdroplet digestion tolerates a high sample injection flow rate, which could reduce the digest collection time. Our study demonstrates that microdroplet could be considered as an emerging, fast and simple protein digestion approach and has the potential for the routine usage in bottom-up protein characterization such as peptide mapping for mAbs.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.2c00072.

Experimental setup of the control experiment, fully tryptic peptides of β-lactoglobulin B, peptide recovery of KVPRNQDWL, nanoESI mass spectra of β-lactoglobulin B, and abundances of semi-specific peptides relative to their corresponding fully tryptic peptides (PDF)

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https://doi.org/10.1021/jacs.2c00072

ACKNOWLEDGMENTS

We thank the National Institutes of Health (1R15GM137311-01) for the financial support of this work.

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