

Structural and functional characterization of solution, gel, and aggregated forms of trypsin in organic solvent-assisted and pH-induced phase changes

Tripsin çözelti, jel ve agregat formlarının organik çözügen içeren ve pH-tektiklenmiş faz geçişlerinde yapısal ve fonksiyonel incelenmesi

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ABSTRACT

Objective: In this study the effect of three different physicochemical parameters on pH-triggered gelation and aggregation of bovine pancreatic trypsin changes and structural and functional changes in these changes in alcohol-water mixtures were studied.

Methods: Trypsin gelation times were studied using inverted tube method. Trypsin stability was studied using trypsin enzyme assay. Protein secondary structural changes were monitored using FTIR spectroscopy. Gel and aggregate macrostructures and morphologies were viewed using Scanning Electron Microscopy.

Results: The solution phase was observed in the absence of both NaOH and CaCl₂. The gel phase was observed in the absence of the either. The aggregate phase was observed in the presence of the both agents all depending on trypsin concentrations used. Trypsin stability studies showed that there were a nearly 53 and 32% specific activity losses after the gelation and aggregation processes. According to FTIR studies β-sheet structure in 1637 cm⁻¹ band disappeared in trypsin gel and trypsin aggregates. Increases in α-helix structure in 1651 cm⁻¹ in trypsin gel and aggregates were observed. Iodoacetamide delayed the gelation and prevented the aggregation indicating the importance of intermolecular disulfides in the both processes.

Conclusion: Trypsin gelation was caused by the denaturation of the protein three dimensional structures. The gel and aggregate formation indicates a secondary structural change towards α-helix structure formation at the expense of β-sheet structure and formation of intermolecular disulfide bonds.

Key Words: Trypsin, Gelation, Aggregation, FTIR

Conflict of Interest: Authors have no conflict of interest.

ÖZET

Amaç: Bu çalışmada alkol-su karışımlarında pH tarafından tetiklenen sığır pankreatik tripsin jelleşmesi ve agregasyonu faz değişiklikleri üzerine üç değişik fizikokimyasal değişkenin yapısal ve fonksiyonel etkileri çalışılmıştır.

Metod: Tripsin jelleşmesi ters tüp metodu ile çalışılmıştır. Tripsin stabilitesi tripsin enzim aktivitesi tayiniyle çalışılmıştır. Protein ikinci yapı değişiklikleri FTIR spektroskopisi metodu ile gözlenmiştir. Jel ve agregat makroyapıları ve morfolojileri taramalı elektron mikroskopisi yöntemiyle incelenmiştir.

Bulgular: NaOH ve CaCl₂ eksikliğinde çözelti fazı gözlenmiştir. Jel fazı bu ikisinden birinin eksikliğinde gözlenmiştir. Agregat fazı ise bu iki ajanın birlikte varlığında tripsin kolantrasyonuna bağlı olarak gözlenmiştir. Tripsin dayanıklılık çalışmaları jelleşmede yaklaşık olarak 53% ve agregasyonda ise 32% özgül aktivitede dayanıklılık kaybı göstermiştir. FTIR spektroskopisi çalışmalarında tripsin jel ve agregatlarında 1637 cm⁻¹ daki β-yaprağı bantının yok olduğu görülmüştür. 1651 cm⁻¹ daki α-heliks bantının güçlenmesi tripsin jel ve agregatlarındaki alfa helix artışına karşılık gelmektedir. Iodoasetamidin jelleşmeyi geciktirmesi jelleşmede moleküller arası disülfid bantlarının önemine işaret etmektedir.

Sonuç: Tripsin jelleşmesinin protein yapısındaki denatürasyondan kaynaklandığı görülmüştür. Jel oluşumu protein ikincil yapısında β-yaprağı kaybına paralel olarak α-heliks oluşumu ve moleküller arası disülfid bağlarının oluşumu ile gerçekleşmektedir.

Anahtar Kelimeler: Tripsin, Jelleşme, Agregasyon, FTIR

Çıkar Çatışması: Yazarların çıkar çatışması yoktur.

Introduction

The fact that proteins are found in different physical forms (solutions, aggregated, gel and powder forms) due to different environmental conditions (pH, ionic strength, or initial protein concentration) has been a very vivid scientific field of biochemistry for years. Due to the well-known structure-function relationship of proteins, function of a protein molecule is linked to environmental conditions. In addition the nature of protein-protein and protein-solvent interactions affect the fate of protein phase behavior.

Under certain conditions some proteins form gels or gel-like structures are commonly recognized as an intermediary phase between solid and solution forms. Gels are regular structures and are composed of network connection of constituting individual particles [1]. Other conditions may also cause aggregation of the protein molecules which have amorphous physical appearance. As opposed to gelation, protein aggregation/precipitation is considered to be a phase separation resulting in formation of a solid phase in a solution [2-3]. The aim of this research is to compare the solution, gel and aggregate forms of trypsin obtained in three different environmental conditions using functional and structural means.

Materials and Methods

Preparation of protein gel and aggregate

Bovine pancreatic trypsin is a globular protein and has a molecular weight of 24.000 daltons. 223 amino acid residues in trypsin are interconnected by six disulphide bonds [4]. Throughout the experiments trypsin was dissolved in 1 mM HCl in the between the concentrations of 2 mg/ml and 10 mg/ml. Then, isopropanol was added to the protein solution at a ratio of 1 (vol/vol). As a last step, NaOH was added to the protein-isopropanol mixture at the designated concentrations. Gelation and aggregation took place depending on the trypsin concentration, CaCl₂ concentration and the pH of the solution.

Inverted tube method

Gelation time was determined when the gel in the tube did not flow or slip through the tube [5].

Trypsin enzyme assay and specific activity

Trypsin catalyzes the hydrolysis of N-benzoyl-L-arginine ethyl ester to N-benzoyl-L-arginine and ethanol. The slope of the absorbance versus time graph is the production rate of BA at pH 7.6 and a total volume of 3.2 ml for the reaction at 25°C. The specific activity of trypsin is equal to (units of activity/ml of solution)/protein concentration (mg/ml). The unit of activity of trypsin activity is defined as 1 µmol BA produced/minute of 0.001 at 253 nm, at pH 7.6 and at 25°C. The reaction was carried out in 100 mM Tris buffer solution with 20 mM CaCl₂ in order to prevent autocatalysis in the conditions of the reaction. Trypsin concentration in solution was determined by UV absorbance at 278 nm.

Iodoacetamide (IAA) studies

Iodoacetamide (IAA) was used as the disulfide blocking agent [6]. 1 M and 2 M IAA were used for 5 mg/ml trypsin concentration to observe the gelation time-aggregation changes.

Scanning Electron Microscopy (SEM) studies

Scanning Electron Microscope (SEM) images were obtained by using Quanta 250 FEG (FEI, USA). SEM images of trypsin, trypsin gel, and trypsin aggregate were taken at 20 µm magnification. All the samples were dried for two days to get rid of the gel humidity. Later the dried samples of the original trypsin, trypsin gel, and trypsin aggregate were exposed to N₂ gas (for coating) and were then analyzed under the SEM.

FT-IR Spectroscopy and spectral analysis

FTIR spectra of trypsin powder, aggregate and gel were recorded with a Perkin-Elmer Spectrum 100 FTIR spectrometer (Perkin-Elmer Inc., Norwalk, CT, USA) equipped with MIR TGS detector. The gel was lyophilized by Labconco FreeZone lyophilizator (Labconco, FreeZone 18 liter freeze dry system) for one night. 225 mg KBr was mixed with 0.5 mg trypsin. The mixture was then exposed to a pressure of 1100 kg/cm² in an evacuated die to produce a KBr pellet for use in FT-IR spectrometer. The spectra were recorded in the 4000–450 cm⁻¹ region at room temperature. A total of 20 scans were taken for each interferogram at 4 cm⁻¹ resolution. Spectrum 100 software (Perkin-Elmer) was used for the data manipulations. From trypsin powder, gel and aggregate (n=3) multiple scans, which gave identical spectra, were performed. These replicates were averaged and the averaged spectra for each sample were then used for further data manipulation and statistical analysis. The spectra were smoothed over 19 points using the Savitzky-Golay algorithm. Then, the spectra were interactively baselined from two arbitrarily selected points. Finally, the spectra were normalized in specific regions for visual comparison of the samples of the amide I bands.

For quantitative analysis of the secondary structures, the gel and aggregate protein secondary structure variations were determined based on intensity calculations from the second derivative spectra of amide I band (1700-1600 cm⁻¹). The spectrum of the second derivation was performed by a Savitzky-Golay algorithm with five smoothing points. The second derivative peaks were normalized. The second derivative signals of the minima peak were used as the peak positions of the original absorption spectra correlation.

Statistical Analysis

The differences between different data groups were compared groups were compared using the Mann-Whitney U Test with the Matlab R2011a program. The statistical results were expressed as means ± standard deviation. A p value of less than 0.05 was considered statistically significant.

Table 1. The effect of iodoacetamide on gelation time and aggregation of trypsin at room temperature.

IAA presence	No IAA	1 M and 2 M IAA
Trypsin gel	2.5 minutes	More than 40 minutes
Trypsin aggregate	Aggregation	No aggregation

Results

The phase behaviour of trypsin as a function of protein concentration, pH and ionic strength

Three different phases were observed due to three different environmental factors: trypsin, CaCl_2 , and NaOH concentrations. Among these environmental factors, NaOH concentration determines the pH of the solution, and CaCl_2 concentration determines the solution's ionic strength. In Figure 1, the phase behavior of trypsin is visualized in three-dimensional space.

In general, gel form was observed between the solution and aggregate forms. Trypsin was shown to be in the solution state at low NaOH and CaCl_2 concentrations. When the physicochemical conditions leading to aggregation and gelation were considered in all trypsin concentrations in the absence of both NaOH and CaCl_2 , solution phase was observed. In the presence of CaCl_2 and in the absence of NaOH, gelation was observed due to binding of CaCl_2 binding to water. In contrast in the presence NaOH and in the absence of CaCl_2 , solution phase was observed due to the insufficient trypsin concentration. In all trypsin concentrations aggregation was observed in the presence of both NaOH and CaCl_2 . Gelation was observed in the presence of either NaOH or CaCl_2 except for the 2 mg/ml trypsin concentration case. This might be because in the presence of NaOH and in the absence of CaCl_2 gelation was not observed due to the insufficient trypsin concentration.

Trypsin stability after gelation/resolubilization

The specific activity measurements were performed to compare the stability of the trypsin following gelation and resolubilization steps. The specific activities of the original trypsin and trypsin gel are shown in Figure 2. The decrease

Table 2. The band assignments of secondary structure sub-bands under Amide I band in 1700-1600 cm^{-1} region [7].

Peak number	Mean frequencies (cm^{-1})	Assignment
1	1694	Antiparallel β -sheets
2	1674	Turns
3	1652	α -helix
4	1643	Random coil
5	1637	β -sheets
6	1625	Aggregated β -sheets

in the specific activity of trypsin was 53.25%, following gelation and resolubilization. The loss in the specific activity indicated irreversible changes in trypsin structure due to phase changes. The decrease in the specific activity of trypsin in the aggregates was 32.45%.

Iodoacetamide (IAA) studies

Iodoacetamide was used as the disulfide bond blocking agent in protein related analyses to investigate the disulfide bond formation in the gelation process as a function of gelation time [6]. In addition, the presence of iodoacetamide prevented aggregation of trypsin. The results indicated that disulfide bonds were important both in the gel and formation processes as shown in Table 1.

Fourier transform infrared spectroscopy studies

FTIR method was applied to observe the secondary structural changes of trypsin which follows gelation and aggregation processes. Amide I band region (between 1600 cm^{-1} and 1700 cm^{-1}) was taken into consideration for protein secondary structure analysis. The band assignments of the original trypsin, trypsin gel, and trypsin aggregate were carried out in the region between 1700-1600 cm^{-1} which corresponded to the amide I region as shown in Figures 3a and 3b. The Amide I band secondary structural assignments of the spectra are shown in Table 2. The changes in the intensities of the bands for the secondary structures are shown in Table 3.

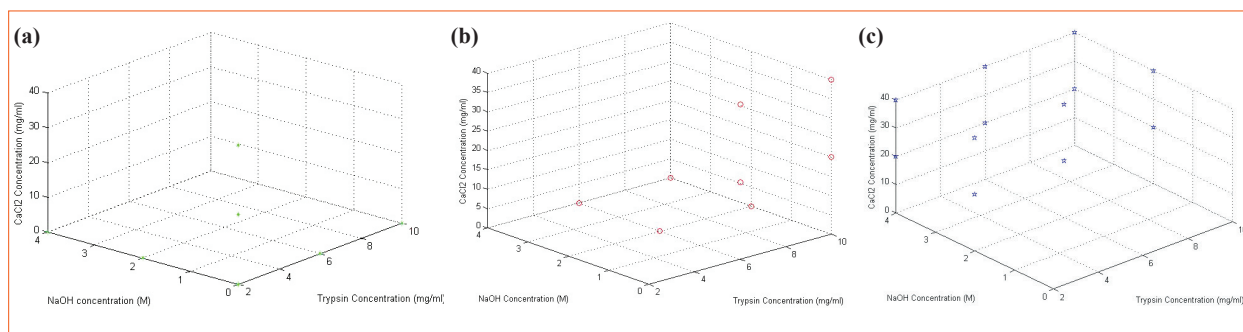


Figure 1. The phase behaviour of trypsin as a function of trypsin concentration, pH and ionic strength in the form of (a) solution (b) gelation, and (c) aggregation.

Table 3. The results of the changes in the intensities of main protein secondary structures for original, gel and aggregated trypsin (Values are the mean \pm standard error of mean for each group).

Functional groups	Trypsin	Gelated trypsin	Aggregated trypsin
Antiparallel β -sheets (at 1694 cm^{-1})	-0.01386 \pm 0.00038	-0.0081 \pm 0.00029	-0.00233 \pm 0.0005
Turns (at 1674 cm^{-1})	-0.01104 \pm 0.00029	-0.00642 \pm 0.00063	-0.00257 \pm 0.00021
α -helix (at 1652 cm^{-1})	-0.01 \pm 0.00011	-0.00497 \pm 0.000367	-0.0045 \pm 0.00056
Random coil (at 1643 cm^{-1})	-0.00978 \pm 0.00063	-0.00798 \pm 0.001	-0.00277 \pm 0.00025
β -sheets (at 1637 cm^{-1})	-0.0167 \pm 0.0003	-0.00565 \pm 0.0003	-0.00253 \pm 0.00035
Aggregated β -sheets (at 1625 cm^{-1})	-0.00956 \pm 0.00009	-0.0069 \pm 0.0011	-0.0026 \pm 0.00026

As seen in Figure 3a, the gel shows changes in secondary structure of trypsin. However, the most significant changes were observed when the second derivative spectra were taken into consideration (Figure 3b). The β -sheets at 1637 cm^{-1} were found to be the ones most affected from the gelation process. The trypsin gel was found to have 32.30% less β -sheet structure ($p < 0.05$). However, the aggregated and anti parallel β -sheet structures were shown to increase by 25.61% and 14.29% in gels respectively ($p < 0.05$). Random coil structure was also observed to increase (36.06%) significantly ($p < 0.05$). There was no percentage change in the percent α -helix structure in the gel structure ($p > 0.05$). Turns structure showed 10.79% increase in the gel phase. The aggregate also showed changes in secondary structure. The aggregate spectra showed a significant band shift in the average spectra compared to the trypsin control spectrum. When the second derivative spectra was taken into consideration, β -sheets which appeared at 1637 cm^{-1} were found to be the ones most affected from aggregation process as seen in Figure 3b. The trypsin aggregate was also found to have 43.52% less β -sheet structure ($p < 0.05$). Similarly the antiparallel β -sheet structure (29.16%) ($p < 0.05$) and turns (8.8%) ($p < 0.05$) were adversely affected by aggregation. However, the aggregated β -sheets were not affected by trypsin aggregation ($p > 0.05$). In contrast to β -sheets, α -helix structure was shown to increase significantly (30.49%) ($p < 0.05$) and

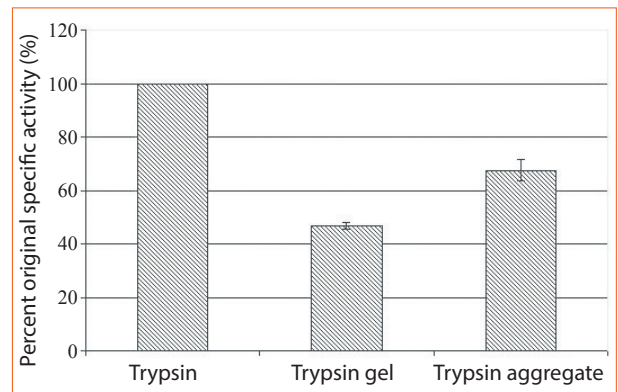


Figure 2. Trypsin stability after gelation and aggregation at 25°C.

random coils increased slightly (2.99%) ($p > 0.05$).

Scanning Electron Microscopy (SEM) studies

The SEM images of trypsin in gel and aggregated phases were analyzed to observe the effect of the gelation and aggregation processes on solid phase trypsin morphology. The SEM images were taken at the magnifications of 20 μm and the results are shown in Figure 4. The original trypsin flakes were observed as thin sheets under the SEM. The SEM images of the gel after drying appeared to have a smoother surface appearance. As seen in the same figure, trypsin aggregates appeared to have an amorphous structure.

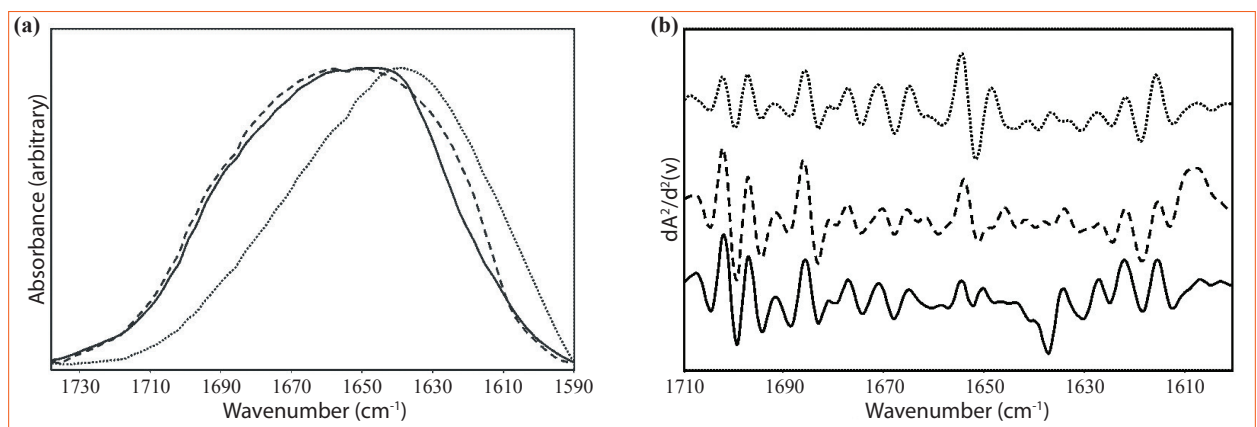


Figure 3. The average (a) absorbance and (b) second derivative spectra of original trypsin, trypsin gel and trypsin aggregate in 1700-1600 cm^{-1} region (continuous line shows original trypsin, dashed line shows trypsin gel, dotted line shows trypsin aggregate).

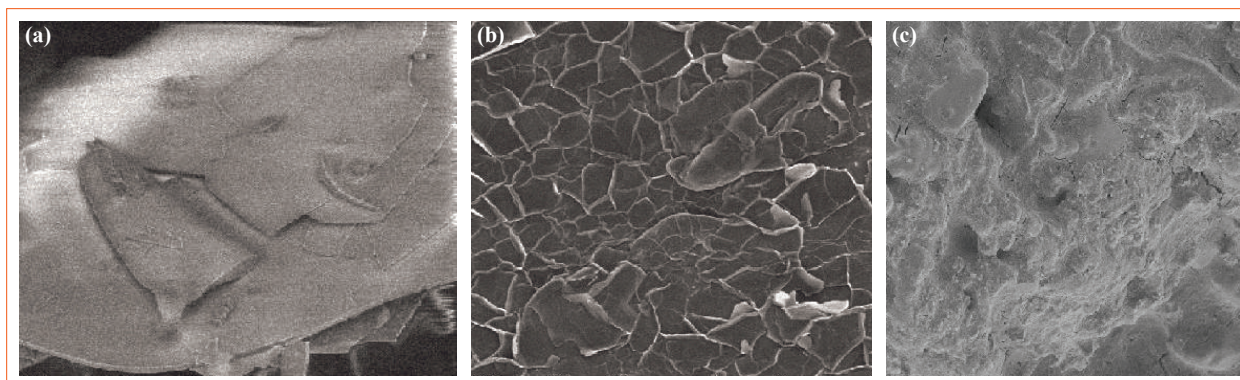


Figure 4. SEM images of (a) original trypsin, (b) trypsin gel, and (c) trypsin aggregate at 20 μm magnifications.

Discussion

In all of the trypsin concentrations studied, the solution phase was observed in the absence of NaOH and CaCl_2 . Gelation was observed in the presence of either CaCl_2 or NaOH. At higher trypsin concentrations (6 mg/ml and 10 mg/ml) gelation was observed because sufficient trypsin concentration for gelation was provided (in the absence of CaCl_2). Aggregation was observed in the presence of both NaOH and CaCl_2 . The results were summarized in a tabular form in Table 4.

The results seem to indicate that at lower CaCl_2 concentrations (2, 6 and 11 mg/ml) CaCl_2 caused neutralization of electrostatic interactions and three dimensional network formations which resulted in gelation. At high CaCl_2 concentrations (22, 33, 44 and 55 mg/ml) where CaCl_2 concentration was in excess for neutralisation, the same charged ions in trypsin molecules started to repel each other and rather than gelation aggregation was observed. Since CaCl_2 binds to water and as the CaCl_2 concentration increased it could hold more water molecules by decreasing the water concentration in protein solution and led to increased protein concentrations simultaneously by increasing the protein-protein interactions. Since CaCl_2 held the distances between water and trypsin molecules shorter, gelation was observed. In cold-set gelation of whey proteins many different salts such as CaCl_2 are commonly used [8,9]. CaCl_2 addition to medium causes neutralization of electrostatic interactions in whey protein gel formation [10]. In a study with whey proteins, it was observed that at low CaCl_2 concentrations (10 mM or

lower than 10 mM), gelation was slow and ordered, fine stranded gels were obtained. At high CaCl_2 concentrations (higher than 10 mM), gelation was fast and aggregates were obtained [11]. At high salt concentrations, electrostatic repulsion between protein molecules screened sufficiently so that protein molecules could come closer to form aggregates [12]. Gelation becomes slower as CaCl_2 concentration reached at low CaCl_2 concentrations as in Kuhn's work [11]. Also Kuhn observed aggregates at high CaCl_2 concentrations.

Protein initial concentration is also an important variable in protein phase changes. Higher trypsin concentrations yielded more interactions between trypsin molecules. As trypsin concentration increased the interaction with the other trypsin molecules, gelation occurred in much shorter times than at lower trypsin concentrations.

Solution pH is another important variable in protein stability and phase behavior [13]. According to Figure 1b, the increasing NaOH concentration leads to a decrease in trypsin solubility and gelation took place. As solubility of trypsin molecules decreased their interaction with each other and environment increased. As a result, a decrease in gelation time was observed. This finding was expected when the basic nature of trypsin is considered ($\text{pI}=10.5$). When the solution pH is near the pI , random aggregation formation is observed. Protein concentration and salt type affects the network formation [14]. According to the experimental observations a model was proposed (Figure 5). Upon triggering of the pH change three different phase behaviors were observed: solution phase, gelation and aggregation. Since different initial conditions led to trypsin

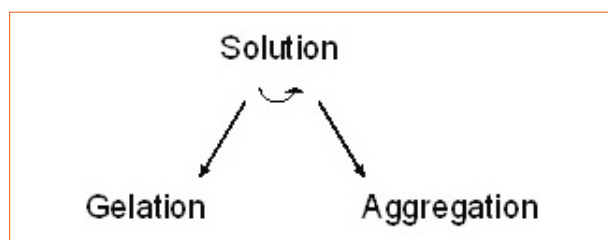


Figure 5. The proposed model for trypsin phase behavior.

Table 4. Trypsin phase behaviour for different CaCl_2 and NaOH concentrations.

NaOH Presence	CaCl_2 Presence	Phase Behaviour
+	+	Aggregation/Gelation
+	-	Solution/Gelation
-	+	Solution/Gelation
-	-	Solution

molecules assumed three different phase behaviors. It was supposed that gelation and aggregation occurred through different molecular mechanisms. This conclusion was further supported by the fact that these different mechanisms resulted in different structural organizations as appeared in the FTIR spectra.

The specific activity results indicated that trypsin partially lost its activity irreversibly after gelation and aggregation. The FTIR spectroscopy results indicated that α -helix structure was found to increase in the aggregation process. This is an unexpected situation since in the most of the protein aggregation studies the amount of α -helix was found to decrease in favor of β -sheet structures [15]. In most of the neurological diseases and other metabolic diseases, β -sheets are the structures leading to the aggregation processes both *in vivo* and *in vitro* conditions [12,16,17,18,19]. As opposed to the aforementioned studies trypsin aggregates were found to have lower β -sheet structures with higher α -helix content in this study.

In our study it was observed that in the presence of IAA, gelation was delayed and also soft gels were formed (Table 1). Since trypsin contains six intramolecular disulfide bonds, the possible roles of intra and inter-molecular disulfide bond formation in the gelation process between trypsin molecules were investigated. Similarly, it is known that disulfide bond formation increases gel hardness. It is well known that protein-protein interactions leading to aggregation are strongly affected by intermolecular β -sheets and disulfide reshuffling. Therefore, intermolecular disulfide bond formation might be responsible for the intermolecular interactions in trypsin gelation and aggregation. Iodoacetamide was found to affect disulfide bonding by thiol blocking. In protein chemistry, it is known that when thiol groups are blocked, disulfide bonding can not take place [6]. In proteins, disulfide bonds are formed by the introduction and removal of the thiol-disulfide exchange reaction [20]. Under adverse physicochemical conditions such as high pressure, temperature, sulfhydryl and cystine groups in proteins are exposed to the surrounding environment and hence facilitate their reaction to form disulfide bonds [20].

Conclusion

Trypsin was shown to form three different phases (solution, gel, and aggregate) for three different physicochemical variables (protein concentration, pH, and ionic strength) in the same experimental system (pH-triggered and alcohol-induced phase change). This system made it possible to investigate the protein structural changes and morphological differences for the three different phases. Contrary to the most of the previous studies, α -helix structures were shown to increase during the phase changes when compared with the solution phase, indicating no contribution from any inter/intra molecular β -sheet structures. Instead, trypsin phase changes were shown to be

driven by inter/intra molecular disulfide trypsin structure.

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Conflict of Interest

There are no conflicts of interest among the authors.

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