Natively Unfolded Regions of the Vertebrate Fibrinogen Molecule

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ABSTRACT Although it has long been realized that a large portion of the fibrinogen α chain has little if any defined structure, the physiological significance of this flexible appendage remains mysterious. Proteins 2006;63:391–397. \odot 2005 Wiley-Liss, Inc.

Key words: fibrinogen; fibrin; intrinsically unfolded domains

INTRODUCTION

In their invitation to this special issue, the Editors noted that authors ought to consider George Rose's research interests in their choice of topic. In line with that request, some explanation may be needed for our chosen subject.

As it happens, in 2002 George Rose edited a volume of Advances in Protein Chemistry on the subject of unfolded proteins.¹ The volume, which contains articles from some of the same individuals contributing to this special issue of Proteins, was comprehensive. Nonetheless, we were surprised to find that a protein that has been the subject of study for a hundred years-and that for more than the last thirty has been thought to have a disproportionate amount of unfolded structure-was never mentioned. Nor was it listed elsewhere in various tallies of protein sequences designated as "intrinsically unstructured" or "natively unfolded."²⁻⁴ The present manuscript is intended to remedy these apparent oversights by reviewing evidence for a lack of structure in certain sectors of the vertebrate fibrinogen molecule, particularly the carboxyl-terminal "domains" of the α chains. At issue are two points: (1) what constitutes a "natively unfolded" entity? And (2) do the fibrinogen α -chain carboxyl-terminal regions meet the requirements? In recent years, criteria for designating a protein as "disordered" or "unfolded"-in whole or parthave usually been based on sensitivity to proteases, on the one hand, or lack of secondary structure as determined by physical methods like circular dichroism, on the other.⁵ Another commonly cited attribute is the absence of interpretable structure in electron density maps calculated from X-ray diffraction data. As we will show, the αC domains of fibrinogen qualify on all these counts and some others as well, including rapid evolutionary change and distinctive amino acid compositions. At the same time, there are some contrary data that imply a minimally folded core structure that need to be discussed also.

STRUCTURAL ASPECTS Vertebrate Fibrinogen

Fibrinogen is the large glycoprotein that circulates in the blood plasma of all vertebrate animals and is the precursor of fibrin blood clots. It is a six-chained entity composed of two pairs each of three polypeptide chains joined by a complex set of disulfide bonds ($\alpha_2\beta_2\gamma_2$). Physicalchemico studies and electron microscopy long ago revealed an extended multi-domained structure approximately 45 nm in length. A plethora of biochemical experiments over the course of half a century gave flesh to that skeletal structure and provided a general if conceptual model of its structure (Fig. 1). One longstanding feature of the model was that the carboxyl two-thirds of the α chains were devoid of secondary structure, oft being referred to as "free-swimming appendages".⁶

The initial basis for this characterization was that the designated segments were easily removed and destroyed by a wide variety of proteases. In the ensuing three decades, an enormous amount of data supporting a mostly unfolded structure has been provided by other means. What follows is a summary of the evidence for that view and some speculation about how inherent flexibility may play a role in fibrin formation.

The Core Fragments D and E

In 1961, a French group digested fibrinogen with the enzyme plasmin and passed the products over an ion exchange column.⁷ Five distinguishable peaks were observed, denoted A–E. Two of these, D and E, accounted for the bulk of the applied material and were destined to become the well studied core fragments D and E. Fragments D, of which there are two per molecule, and E (the central domain) together account for two-thirds of the mass of the starting fibrinogen. The missing one-third, which in most mammalian molecules amounts to about a thousand residues per native dimer, is mostly from the carboxyl-terminal segments of the two α chains. Fragments D and E also accounted for the main morphological features observed in shadow-cast electron micrographs of fibrinogen.^{8,9}

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Fig. 1. An old model of fibrinogen concocted on the basis of early electron microscopy studies and sundry biochemical data. FPA, fibrinopeptide A; FPB, fibrinopeptide B; SS RING, disulfide rings; XL, cross-linking sites; CHO, carbohydrate. A α , β , and γ denote the α , β , and γ chains of the dimeric protein, respectively. Note the minimally folded α C regions. (reprinted from Marder, Francis and Doolittle⁷¹)



Fig. 2. Diagram of various regions of human fibrinogen α chain. Residue numbers denote some key points. A proteolytic fragment⁴³ and two recombinantly expressed regions^{28,56} are labeled by authors and year. Positions corresponding to changes that lead to truncated α chains in some variant human fibrinogens are marked "X" (fibrinogens Arnhem⁴⁸ and Marburg⁵¹). Positions at which changes lead to amyloid formation are denoted by shaded circles.

Moreover, when fibrinogen was subjected to limited proteolysis, the percent of secondary structure remaining in the core structure, as determined by circular dichroism, increased greatly.¹⁰ This observation was consistent with the notion that the three globules observed in electron micrographs were likely connected by α -helical "coiled coils."¹¹ Conversely, CD studies on the material released gave a signal characteristic of a "random coil".^{12,13}

On another front, elegant scanning calorimetry experiments showed that there were two discrete moieties in fibrinogen that "melted" at very different temperatures, two widely separated heat-absorbing peaks being entirely accounted for by the fragments D and E with no other obvious features being evident.¹⁴

Amino Acid Sequence Studies

When the human fibrinogen α chain was sequenced,¹⁵ it was found to have three quite characteristic zones: a coiled-coil domain that was homologous to the β and γ

chains, a region consisting of ten 13-residue imperfect repeats, and an approximately 250-residue long terminal domain corresponding to what is now generally referred to as the αC domain (Fig. 2). The sequence was wholly consistent with the model shown in Figure 1 that had a terminal αC region connected to the main-frame of the molecule by a flexible tether composed of imperfect repeats, both sections being susceptible to release by proteases.

Sequence studies on other species revealed that the carboxyl-terminal regions of α chains have experienced a very rapid rate of evolutionary change, not only with regard to amino acid substitution, but also in the way of numerous deletions and insertions.¹⁶ A region containing a single disulfide bond is modestly conserved (Fig. 3), but even it changes faster than the average plasma protein. For example, the 40-residue conserved segment containing the disulfide is only 67% identical in humans and chickens, whereas the entire 86-kDa fragments D of these two



Fig. 3. Schematic depiction of human, bovine, rat and chicken fibrinogen α chains. The regions with the variable numbers of repeats and the α C domains are both readily removed by limited proteolysis. Note that the chicken α chain lacks the repeat region which in other species provides a connecting tether for the terminal α C domain.

Human	GSFRPDSP	GS G	NARE	NNPDWG	TF	EE	VSGNVS	PGT RRE	YH TEKLVTSKGDK
Rat	GSSRP SS	GS G	NLKE	SNPDWG	EF	SE	FGGSSS	PAT RKE	YH T GK LVTSKGDK
Cow	SSFRPDSS	GH G	NIRP	SSPDWG	TF	RE	EGSVS	SGT KQE	FH T GK LVTTKGDK
Chicken	PSPEMQAM	SAFN	NIKO	VOM	VL	ERPET	DHVAEA	RGD SSP	SH TGKLITSSHRR
Xenopus	KDINOYSF	VLEG	337253.04		TK	EE	VKGS S	IGT SSV	VFPSGSGKDVDFOGSKG
Zebrafish	LDKOMDNL	NAMR	VOSVE	TVSSLG	IMKSRPL	KD VL	VPTIYK	SGTGKAEOK	LLFGDVGOMOLSLEAEG
							*		
Human	ELRTGKEK	V TS	GST	TTTRRS	CSKTVT	K TVI	GPDG H	KEVTKEVVT	S EDGSDC P
Rat	ELLIGNEK	V TS	TGT	STTRRS	CSKTIT	K TVL	GNDG H	REVVKEVVT	S DDGSDC G
Cow	ELLIDNEK	V TS	GHT	TTTRRS	CSKVIT	K TVT	NADG R	TETTKEVVK	S EDGSDC G
Chicken	ESPSLVDK	T SS	A	SSVHR	CTRTVT	KKVIS	GPDGPR	EEIVEKMVS	S DGSDC
Xenopus	TOSGTVKGGDO	S TK	VIT	ESRTVS	CTKTKK	KRVII	TPEG P	KE EIIE	SYSGGAECEKLGKFSTG
Zebrafish	STAESAATVSK	FPTS	GTGST	SQTSSK	CTKSVR	KVITH	TKDGPV	EKV EVSS	S GPGC D
Human	EAMDLG	TL	SGIG	TLDGFR	HRHPDEA	AFF	DTASTG	KTFPGFFSP	MLGEFVSETESRGSE S
Rat	DGMDLGL	тн	SFSG	RLDELSI	RMHPELG	SFY	D	SRF GSLTS	NFKEFGSKT SD S
Cow	DA DEDW	HHT	FPSRG	NLDDFFI	HRDKD			DFFTR	SSHEFDGRT GL A
Chicken	SHLQGGREG	STYH	FSGTO	DFHKLDI	RLLPDLE	SFFTH	DSVSTS	SRHSIGSST	SSHVTGAGSSHLGTGGK
Xenopus	DTFEAGSDGTR	VIK	ISGSS	SLGDIS	KIPSFE	EFL	SSSGR	KTQTSQTSS	SSSSSSSSTYSKGVT H
Zebrafish	DLGKLGIS	DSEF	LTAAB	DGKDLSI	R DGL	KI	TVTGGD	EKSLTSLSH	STSDLGG FGGD
	*								
Human	GIFTNTKES	SSH	HPGI	AEF	PSRGK	SSSY	SKQ F	TSSTSYNRG	DSTFESKSY K MADEAGS
Rat	DIFTDIENP	SSH	V	PEF	SSSSK	TSTV	RKQ V	TKS	YKMADEAAS
Cow	PEFAALGES	GS			SSSK	TSTH	SKQFV	SSSTTVNRG	GSAIESKHF K MEDEAES
Chicken	DKFTDLGEEE	EDD	FGGLQ	PSGF	AAG	SASH	SKTVLT	SSSSSFNKG	GSTFETKSL K TRETSEQ
Xenopus	DAFSDLGEDEF	DDFS	HTGLD	SPSF	TPIQKE	GSSTY	TKTVVS	SSSRGSTKD	GTQFDIKSM K SGPVFDD
Zebrafish	FFKNLGTDNR	KYAS	SSSST	STKTVSI	DGSKSYS	KTTLT	SDPIFG	DDLGAFMRG	DVEEDMPDIHARSVKS
Human	EADHEG THS	TKRG	HAKSF	RPV					
Rat	EAHQEGDTRT	TKRG	RARTM	IRGIHA					
Cow	LEDLGFKGAHG	TQKG	HTKAP	PRA					
Chicken	LGGVQHDQSAE	DTPD	FKARS	FRPA					
Xenopus	FGPIQHDNSEE	DRPD	FQART	VRKE					
Zebrafish	QDDRKAG	FVGG	GTLE.						

Fig. 4. Sequence comparisons of α C domains from six assorted vertebrates. The extensive differences reflect the rapid evolutionary change that occurs in this region of the fibrinogen molecule. Only six residues in the approximately 240-residue segment are absolutely conserved (denoted by asterisks). Those positions in which five of the six residues are identical are emboldened. The most conserved region, delineated by the two cysteine residues that form the single disulfide bond, is shaded.

species are almost 80% identical. In general, the amino acid compositions of the α C domains in various species their rapid change not withstanding—are consistent with what is now being deigned as characteristic of intrinsically or natively unfolded proteins.^{17,18} Rapid evolution has been reported for other proteins with long disordered regions.¹⁹ As it happens, also, some intrinsically disordered domains have been shown to be the result of evolutionary expansion of repeat regions.²⁰ In this regard, it is significant that the number of repeats in the central region of these chains varies widely from species to species (Fig. 4). Indeed, even the sizes of the repeats differ, being 13 residues in most species, but 15 residues in marsupials, and 18 residues in dogs¹⁶ and lampreys.²¹ Birds were found to lack the repeated region completely,²² as has now been found to be the case for bony fish and amphibians.²³ It has been proposed that in those molecules that do have the repeats they serve as an accordion or spring-like tether for the α C domain.²¹

Evidence Favoring a Compact aC Domain.

In calorimetry studies conducted subsequent to the those cited above, bovine fibrinogen was found to exhibit additional small but reproducible heat effects that were consistent with some minimal structure in the α C domains. These data were regarded as indicative of a compact domain connected to the main body by a flexible tether.^{24,25} Subsequent studies on human fibrinogen did not find the same thermal effects, however.²⁶ Nonetheless, the view has persisted that these are *compact globular* domains.²⁷ The question arises as to whether the descriptors "compact" and "globular" imply a fixed and folded structure. Although "globular" is a broad, all-embracing term that can accommodate both folded and unfolded domains, "compact" almost always implies a degree of order.

In line with this view, more recent studies employing fluorescence methods, differential scanning calorimetry, and CD were conducted on recombinant constructs of both the bovine and human αC domain, with and without the connecting tethers²⁸ (Fig. 2). Although thermally-induced changes in ellipticity were detected, virtually no conventional secondary structure (i.e., α -helix or β -structure) was found. Instead, the evidence favored the existence of an extended helical poly(L-proline) type II structure, mostly associated with the connector (tether) region. It is notable that the polyproline II helix is now regarded as a hallmark of the unfolded state.^{1,29} It might be noted, also, that not all α -chain repeats in different species are proline-rich; but then again, polyproline II structure is often observed in other proline-poor proteins, as well.³⁰ As pointed out to us by a reviewer, enthalpy changes during scanning calorimetry are given to various interpretations and must be considered with great caution.³¹

Electron Microscopy

There have been conflicting reports over the years about the whereabouts of the α C domain as determined by various electron microscope approaches, some of which contend that the α C domains are associated with the central domain,^{32,33} and others, mostly utilizing antibodies (or FAB fragments) directed against peptides in the α C region, which find the immuno-reactive material in varied and more distal locations.^{34,35} In the latter experiments, only the antibodies (or FAB fragments), and not the α C regions themselves, appeared to be shadowed. However, there have been more definitive studies on isolated α C domains, particularly focused on a 40-kD fragment from bovine fibrinogen.³⁶

A recent report utilizing scanning-transmission electron microscopy (STEM) has focused on a minor form of fibrinogen that has alternatively spliced α chains with carboxylterminal globular domains (denoted α C') homologous to the β and γ chains located distal to (in a sequence sense) the usual α C domains.³⁷ In those micrographs allegedly including both kinds of domain, the globular α C' domains, which contain about the same number of residues as α C domains, dwarf the images of any neighboring material.

Crystal Structures

More than three decades ago, it was serendipitously found possible to crystallize bovine fibrinogen that had been trimmed down by limited proteolysis, even though it was not possible to crystallize the native protein.³⁸ Eventually, a structure of this modified protein—lacking αC domains—was determined at a nominal 4 Å resolution.³⁹

The fact that chicken fibrinogen totally lacks the middleregion tether²² (Fig. 4) prompted efforts to crystallize that protein. Indeed, the native protein was crystallized and a structure determined at 2.7 Å resolution.⁴⁰ Not surprisingly, the α C domains were not discernable. It is indisputable that the α C domains in these crystalline settings are moving, whether or not they are intrinsically unfolded, and this in a fibrinogen whose α chains lack the repetitive region thought to be the flexible tether. Efforts to crystallize recombinant constructs corresponding to the α C domains from both bovine and chicken fibrinogens have been wholly unsuccessful.⁴¹

All of these observations are in accord with the αC domains of fibrinogen having the properties of "natively unfolded proteins" as currently defined. The questions arise, why are these regions of fibrinogen unfolded, and what is their raison d'etre?

FUNCTIONAL ASPECTS

Fibrin Formation

There is general agreement that αC domains play a role in fibrin formation. Antibodies directed to that region of the molecule interfere with clotting,⁴² as do purified fragments themselves.^{36,43,44} Moreover, if the αC domains are absent, whether removed by proteases or—as we will discuss further below—the result of genetic aberration, clotting is greatly slowed.⁴⁵ What the role may be, however, is not at all clear.

Put very briefly, fibrin formation occurs in two stages. In the first, thrombin removes the fibrinopeptides A from the amino-terminal segments of α chains, thereby exposing a set of knobs in the central region of fibrinogen that can fit into holes at the extremeties of other molecules and bridge them together in an overlapping fashion to form a twomolecule thick protofibril. In the second stage, thrombin removes the fibrinopeptide B from the amino-terminal segment of the β chain, the result of which is the lateral association of the protofibrils and the formation of mature fibers. The involvement of the readily removed α C domain appears to be limited to this second stage.

In this regard, it has long been known that fibrinogen prepared by conventional means from normal plasma contains separable fractions that have different degrees of solubility.⁴⁶ Comparisons of "high-soluble" and "low-soluble" fractions have been traced to differences in the carboxyl-terminal regions of the α chains, the more soluble fractions having distinctly shorter chains, the result of in vivo proteolysis.⁴⁷ Similar results are observed when limited proteolysis is performed in vitro.

As it happens, however, proteolysis—especially when conducted in vitro—removes other parts of the fibrinogen molecule, most notably a flexible 40–50 residue segment from the amino-terminus of the β chain, and the possible contribution of these missing segments to the slowed clotting could not be discounted, even though in some reports great care was taken to show that the β chains

Genetically Defective Fibrinogens

Genetically variant forms of human fibrinogen have been helpful in assessing the functions of the αC domain. A number of different variants result in truncated α chains. the most informative of which are arguably those that occur in the region just before the beginning of the ten imperfect repeats and which ought to be equivalent to those generated by limited proteolysis in vitro. One of these is fibrinogen Arnem,⁴⁸ in which a residue ordinarily cleaved by plasmin, aLys-219, has its codon mutated to a terminator. The patient experienced bleeding problems, and the fibrinogen was definitely slow to clot when treated with thrombin. Similar situations occur in a host of other variants in which the α -chain changes are more distal. On their own, then, variant human fibrinogens lacking all or part of the αC domain demonstrate the importance of that structure in fibrin formation.

Some of the genetically variant fibrinogens provide evidence for the flexible character of αC domains, as well as providing circumstantial evidence about their unfolded nature. In this regard, a variety of changes lead to the presence of unpaired cysteine residues in the α C region. These come about in two ways. In one these, noncysteine residues are simply mutated to cysteines. An example is Fibrinogen Dusart, which has a substitution of a cysteine for an arginine at position α 554.^{49,50} In the other situation. termination codons occur in the 29-residue segment between cysteines $\alpha 442$ and $\alpha 472$ (Fig. 2). An example is fibrinogen Marburg which has a stop codon in place of the normal lysine at position $\alpha 461$,⁵¹ and as a result it not only lacks the carboxyl-terminal 150 amino acids that encompass most of the αC domain but has an unpaired cysteine at $\alpha 442$ as well.

Invariably these unpaired cysteines form disulfide bonds with plasma albumin, the most abundant of the plasma proteins. Burdened by that extra 67-kDa entity, these fibrinogens exhibit markedly abnormal polymerization. Significantly, none has ever been found to be disulfide-linked to its partner α C domain in the same or other molecules. It is of interest, also, that electron microscopy readily identifies the attached plasma albumin moieties but not the α C domains themselves.⁵²

It should be mentioned in passing that several variant human fibrinogens in the region of α -chain residues 522–554 result in the formation of amyloid deposits.^{53–55} Interestingly, unfolded domains are often associated with amyloid formation.² Finally, adding another note to the puzzle is the observation that a recombinant fibrinogen engineered to have its α chains terminate at position α 251 was only modestly affected with regard to polymerization.⁵⁶

Factor XIII Cross-Linking

In its early stages, fibrin formation is driven by the formation of noncovalent interactions. As the molecules polymerize and specific regions are brought into juxtaposition, the transglutaminase known as factor XIIIa incorporates isopeptide linkages between certain lysine and glutamine side chains. In particular, sets of reciprocal crosslinks are rapidly formed between the carboxyl-terminal segments of abutting γ chains—in a time course measured in seconds or minutes-and while the first stage of fibrin formation is still in progress.^{57,6} In contrast, a network of the same kind of cross-links (ϵ -amino- γ -glutamyl lysines) is formed very much more slowly-over the course of hours or days—between αC domains. The latter involve a variety of lysine and glutamine side-chains⁵⁸ and lead to a multimerically cross-linked system.⁵⁹ Studies with recombinant αC domains and exogenous factor XIII have yielded similar results.^{60,61}

Unquestionably, the cross-linking of the αC domains contributes to the stability of clots, even though the wide variety of different cross-links formed reflects an almost random nature of encounter consistent with a lack of specific structure.⁶²

The αC Domain as a Binding Site

It is well established that the α C domains in fibrinogen have binding sites for factors that are involved in the destruction of fibrin clots, namely, tissue plasminogen activator (t-PA) and plasminogen.⁶³ It is also known that inhibitors of such proteolytic enzymes can bind to these regions.⁶⁴ Moreover, these same regions of the α chains are also known to be involved in the binding of fibrinogen to blood platelets,⁶⁵ fibroblasts,⁶⁶ endothelial,⁶⁷ and other lymphoid cells.⁶⁸ The α C domain is also the site for cross-linking fibrinogen to fibronectin,⁶⁹ a process vital to wound healing.

A Guardian Function

A region of a protein can have multiple functions, of course. One interesting but almost forgotten suggestion was that these flexible appendages—like leashed hounds—guard the otherwise vulnerable coiled coils against proteolytic attack. A careful kinetic analysis of the proteolysis of both bovine and human fibrinogens concluded that such a role was indeed reasonable.⁷⁰

SUMMARY

In spite of the evidence for an intrinsically disordered structure marshaled above, the popular view remains that α C domains are compact and globular.^{27,28} That in most species they are tethered to the main frame of the parent molecule by a flexible connector is not in dispute. How these regions can be associated *intra*-molecularly in fibrinogen and be in motion at the same time remains mysterious. It may be that these are matters of degree: we need to ask, how disordered and how much motion?

Beyond that, the argument may be largely semantic, different images being provoked by the terms "compact globular" on the one hand, and "intrinsically unfolded" on

the other. In this regard, reference is made in discussions of other natively unfolded proteins to "an ensemble of interchanging conformations."¹⁸ Surely this must be the case for the αC domain, defined loops of as yet unknown conformations being able to serve as specific binding sites for other proteins. Other loops must be able to form loose associations with themselves or other features on the neighboring αC domains. What these domains do not do is form stable structures with defined geometries that form specific associations with themselves; if they did we would expect readily defined cross-links to be introduced by factor XIII, if only because of the very limited number of glutamines that can act as acceptors. We would also expect to see the *intra*-molecularly associated structures in X-ray diffraction patterns. Finally, we would expect regions of association-whether intra- or inter-molecularly-to have amino acid sequences that are at least moderately conserved in an evolutionary sense.

In the end, there is good reason to believe that the αC domains of vertebrate fibrinogen are intrinsically unfolded. Even so, their mobility must be greatly constrained after the conversion of fibrinogen to fibrin, and especially after the introduction of covalent cross-links between them. The fact that individual cross-links involve different glutamine and lysine participants underscores the randomized encounters expected from these highly mobile entities. Apart from this ultimate stabilization, the general whereabouts and precise function of these segments remain controversial matters.

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REFERENCES

- 1. Rose GD. Getting to know u. Adv Prot Chem 2002;62:xv-xxi.
- Wright PE, Dyson HJ. Intrinsically unstructured proteins: reassessing the protein structure-function paradigm. J Mol Biol 1999;293:321–331.
- Uversky VN, Gillespie JR, Fink AL. Why are "natively unfolded" proteins unstructured under physiologic conditions? Proteins 2000; 41:415–427.
- Dunker AK, Brown CJ, Lawson JD, Iakoucheva LM, Obradovic Z. Intrinsic disorder and protein function. Biochemistry 2002;41: 6573–6582.
- Garner E, Cannon P, Romero P, Obradovic Z, Dunker AK. Predicting disordered regions from amino acid sequence: common themes despite differing structural characterization. Genome Inform 1998;9:201–213.
- Doolittle RF. Structural aspects of the fibrinogen to fibrin conversion. Adv Protein Chem 1973;27:1–109.
- Nussenzweig V, Seligmann M, Pelimont J, Grabar P. Les produits de degradation du fibrinogene humain par la plasmine. Annal L'institut Pasteur 1971;100:377–389.
- 8. Hall CE, Slayter HS. The fibrinogen molecule: its size, shape and mode of polymerization. J Biophys Biochem Cytol 1959;5:11–17.
- Fowler WE, Fretto LJ, Erickson HP, McKee PA. Electron microscopy of plasmic fragments of human fibrinogen as related to trinodular structure of the intact molecule. J Clin Invest 1980;66: 50–56.
- Huseby RM, Mosesson MW, Murray M. Studies of the amino acid conformation of human fibrinogen: comparison of fractions I-4 and I-8. Physiol Chem Physics 1970;2:374–384.
- Cohen C. Invited discussion at 1960 Symposium on Protein Structure. J Pol Sci 1961;49:144-145.
- 12. Budzynski AZ. Difference in conformation of fibrinogen degrada-

tion products as revealed by hydrogen exchange and spectropolarimetry. Biochim Biophys Acta 1971;229:663-671.

- 13. Takagi T, Doolittle RF. Amino acid sequence on the α -chain of human fibrinogen: location of four plasmin attack points and a covalent crosslinking site. Biochemistry 1975;14:5149–5156.
- Donovan JW, Mihalyi E. Conformation of fibrinogen: calorimetric evidence for a three-nodular structure. Proc Natl Acad Sci USA 1974;71:4125-4128.
- 15. Doolittle RF, Watt KWK, Cottrell BA, Strong DD, Riley M. The amino acid sequence of the α -chain of human fibrinogen. Nature 1979;280:464-468.
- Murakawa M, Okamura T, Kamura T, Shibuya T, Harada M, Niho Y. Diversity of primary structures of the carboxy-terminal regions of mammalian fibrinogen Aα-chains. Thromb Haemostasis 1993;69:351–360.
- Bracken C, Iakoucheve LM, Romero PR, Dunker AK. Combining prediction, computation and experiment for the characterization of protein disorder. Curr Opin Struct Biol 2004;14:570-576.
- Oldfield CJ, Cheng Y, Cortese MS, Brown CJ, Uversky VN, Dunker AK. Comparing and combining predictors of mostly disordered proteins. Biochemistry 2005;44:1989–2000.
- Tompa P. Intrinsically unstructured proteins evolve by repeat expansion. BioEssays 2003;25:847-855.
- Brown CJ, Takayana S, Campen AM, Vise P, Marshall TW, Oldfield CJ, Williams CJ, Dunker AK. Evolutionary rate heterogeneity in proteins with long disordered regions. J Mol Evol 2002;55: 104–110
- Doolittle RF. The structure and evolution of vertebrate evolution: a comparison of the lamprey and mammalian proteins. Adv Exp Med Biol 1990;281:25–37.
- 22. Weissbach L, Grieninger G. Bipartite mRNA for chicken α -fibrinogen potentially encodes an amino acid sequence homologous to β and γ -fibrinogens. Proc Natl Acad Sci USA 1990;87:5198–5202.
- Benson DA, Karsch-Mizrachi I, Lipman DJ, Ostell J, Wheeler DL. GenBank. Nucl Acids Res 2005;33:D34–D38.
- Privalov PL, Medved LV. Domains in the fibrinogen molecule. J Mol Biol 1982;159:665–683.
- 25. Medved LV, Gorkun OV, Privalov P. Structural organization of C-terminal parts of fibrinogen A α -chains. FEBS Lett 1983;160:291–295.
- Procyk R, Medved L, Engelke KJ, Kudryk B, Blomback B. Nonclottable fibrin obtained from partially reduced fibrinogen: characterization and tissue plasminogen activator stimulation. Biochemistry 1992;31:2273–2278.
- 27. Weisel JW, Medved, L. The structure and function of the αC domains of fibrinogen. Ann NY Acad Sci 2001;936:312–327.
- 28. Tsurupa G, Tsonev L, Medved L. Structural organization of the fibrin(ogen) αC domain. Biochemistry 2002;41:6449–6459.
- Stapley BJ, Creamer TP. A survey of left-handed polyproline II helices. Protein Sci 1999;8:587–595.
- Cao W, Bracken C, Kallenbach NR, Lu M. Helix formation and the unfolded state of a 52-residue helical protein. Protein Sci 2004;13: 177–189.
- Richardson JM, Makhatadze GI. Temperature dependence of the thermodynamics of helix-coil transition. J Mol Biol 2003;335:1029– 1037.
- Mosesson MW, Hainfeld J, Wall J, Haschemeyer RH. Identification and mass analysis of human fibrinogen molecules and their domains by scanning transmission electron microscopy. J Mol Biol 1981;153:695–718.
- Erickson HP, Fowler WE. Electron microscopy of fibrinogen, its plasmic fragments and small polymers. Ann NY Acad Sci 1983;408: 146–163.
- Telford JN, Nagy JA, Hatcher PA, Scheraga HA. Location of peptide fragments in the fibrinogen molecule by immunoelectron microscopy. Proc Natl Acad Sci USA 1980;77:2372–2376.
- Price TM, Strong DD, Rudee ML, Doolittle RF. Shadow-cast electron microscopy of fibrinogen with antibody fragments bound to specific regions. Proc Natl Acad Sci USA 1981;78:200–204.
- 36. Veklich YI, Gorkun OV, Medved LV, Nieuwenhuizen W, Weisel JW. Carboxyl-terminal portions of the alpha chains of fibrinogen and fibrin. Localization by electron microscopy and the effects of isolated alpha C fragments on polymerization. J Biol Chem 1993;268:13577-13585.
- Mosesson MW, DiOrio JP, Hernandez I, Hainfeld JF, Wall JS, Grieninger G. The ultrastructure of fibrinogen-420 and the fibrin-420 clot. Biophys Chem 2004;112:209-214.

- Tooney N, Cohen C. Microcrystals of a modified fibrinogen. Nature 1972;237:23–25.
- Brown JH, Volkmann N, Jun G, Henschen-Edman AH, Cohen C. Crystal structure of a modified bovine fibrinogen. Proc Natl Acad Sci USA 2000;97:85–90.
- Yang Z, Kollman JM, Pandi L, Doolittle RF. Crystal structure of native chicken fibrinogen at 2.7 Å resolution. Biochemistry 2001; 40:12515–12523.
- Kollman J. Structural studies of the *Panulirus interruptus* clottable protein [dissertation]. San Diego: University of California; May, 2005.
- 42. Cierniewski CS, Plow EF, Edgington TS. Conformation of the carboxy-terminal region of the A α chain of fibrinogen as elucidated by immunochemical analyses. Eur J Biochem 1984;141:489–496.
- 43. Lau HKF. Anticoagulant function of a 24-Kd fragment isolated from human fibrinogen A α chains. Blood 1993;81:3277–3284.
- Gorkun OV, Veklich YI, Medved LV, Henschen AH, Weisel JW. Role of the alpha C domains of fibrin in clot formation. Biochemistry 1994;33:6986-6997.
- Medved LV, Gorkun OV, Manyakov VF, Belitser VA. The role of fibrinogen αC domains in the fibrin assembly process. FEBS Lett 1985;181:109-112.
- 46. Mosesson MW, Alkjaersig N, Sweet B, Sherry S. Human fibrinogen of relatively high solubility. Comparative biophysical, biochemical, and biological studies with fibrinogen of lower solubility. Biochemistry 1967;6:3279–3287.
- 47. Holm B, Brosstad F, Kierulf P, Godal HC. Polymerization properties of two normally circulating fibrinogens, HMW and LMW. Evidence that the COOH-terminal end of the α -chain is of importance for fibrin polymerization. Thromb Res 1985;39:595– 606.
- Vleitman JJ, Verhage J, Vos HL, van Wijk R, Remijn JA, van Solinge WW, Brus F. Congenital afibrinogenaemia in a newborn infant due to a novel mutation in the fibrinogen Aalpha gene. Br J Haematol 2002;119:282–283.
- 49. Koopman J, Haverkate F, Grimbergen J, et al. The molecular basis for fibrinogen Dusart (A α 554 Arg > Cys) and its association with abnormal fibrin polymerization. J Clin Invest 1993;91:1637–1643.
- Collet JP, Woodhead JL, Soria J, Soria C, Mirshahi M, Caen JP, Weisel JW. Fibrinogen Dusart: electron microscopy of molecules, fibers and clots, and viscoelastic properties of lots. Biophys J 1996;70:500-510.
- Koopman J, Haverkate F, Grimbergen J, Egbring R, Lord ST. Fibrinogen Marburg: a homozygous case of dysfibrinogenemia, lacking amino acids Aα 461-610 (Lys 461 AAA > Stop TAA). Blood 1992;80:1972–1979.
- 52. Siebenlist KR, Mosesson MW, DiOrio JP, Soria J, Soria C, Caen JP. The polymerization of fibrinogen Dusart (Aα554 Arg > Cys) after removal of carboxy terminal regions of Aα-chains. Blood Coag Fibrinolysis 1993;4:61-65.
- 53. Benson MD, Liepnieks J, Uemichi T, Wheeler G, Correa R. Hereditary renal amyloidosis associated with a mutant fibrinogen α -chain. Nature Gen 1993;3:252–255.
- 54. Uemichi T, Liepnieks J, Benson MD. Hereditary renal amyloid-

osis with a novel variant fibrinogen. J Clin Invest 1994;93:731–736.

- 55. Uemichi T, Liepnieks J, Yamada T, Gertz MA, Bang N, Benson MD. A frame shift mutation in the fibrinogen $A\alpha$ chain gene in a kindred with renal amyloidosis. Blood 1996;87:4197–4203.
- 56. Gorkun OV, Henschen-Edman AH, Ping LF, Lord ST. Analysis of Aα251 fibrinogen: the αC domain has a role in polymerization albeit more subtle than anticipated from the analogous proteolytic fragment X. Biochemistry 1998;37:15434-15441.
- Chen R, Doolittle RF. γ-γ Cross-linking sites in human and bovine fibrin. Biochemistry 1970;10:4486-4491.
- 58. Sobel JH, Gawinowicz MA. Identification of the α chain lysine donor sites involved in factor XIIIa fibrin cross-linking. J Biol Chem 1996;271:19288–19297.
- McKee PA, Mattock P, and Hill RL. Subunit structure of human fibrinogen, soluble fibrin and cross-linked insoluble fibrin. Proc Natl Acad Sci USA 1970;66:738–744.
- Matsuka YV, Medved LV, Migliorini MM, Ingham KC. Factor XIIIa-catalyzed cross-linking of the recombinant αC fragments of human fibrinogen. Biochemistry 1996;35:5810–5816.
- Tsurupa G, Veklich Y, Hantgan R, Belkin AM, Weisel JW, Medved L. Do the isolated fibrinogen αC-domains form ordered oligomers? Biophys Chem 2004;112:257–266.
- 62. Yang Z, Mochalkin I, Doolittle RF. A model of fibrin formation based on crystal structures of fibrinogen and fibrin fragments complexed with synthetic peptides. Proc Natl Acad Sci USA 2000;97:14156-14161.
- Tsurupa G, Medved L. Identification and characterization of novel t-PA- and plasminogen-binding sites within fibrin(ogen) αCdomains. Biochemistry 2001;40:801–808.
- 64. Kimura S, Aoki N. Cross-linking site in fibrinogen for α 2-plasmin inhibitor. J Biol Chem 1986;261:15591–15595.
- 65. Hawiger J, Kloczewiak M, Bednarek MA, Timmons S. Platelet receptor recognition domains on the α chain of human fibrinogen: structure-function analysis. Biochemistry 1989;28:2909–2914.
- Farrell DH, Al-Mondhiry HA. Human fibroblast adhesion to fibrinogen. Biochemistry 1997;36:1123–1128.
- Cheresh DA, Berliner SA, Vicente V, Ruggeri ZM. Recognition of distinct adhesive sites on fibrinogen by related integrins on platelets and endothelial cells. Cell 1989;58:945–953.
- Plow EF, Edgington TS. Lymphocyte suppressive peptides from fibrinogen are derived predominantly from the Aα chain. J Immunol 1986;137:1910-1915.
- Makogonenko E, Tsurupa G, Ingham K, Medved, L. Interaction of fibrin(ogen) with fibronectin; further characterization and localization of the fibronectin-binding site. Biochemistry 2002;41:7907– 7913.
- Shrager RI, Mihalyi E, Towne DW. Proteolytic fragmentation of fibrinogen. II. Kinetic modeling of the digestion of human and bovine fibrinogen by plasmin and trypsin. Biochemistry 1976;15: 5382-5386.
- Marder VJ, Francis CW, Doolittle RF. In: Colman RW, Hirsh J, Marder VJ, Salzman EW, editors. Fibrinogen structures and physiology in Hemostasis and Thrombosis: Basic Principles and Clinical Practice. Philadelphia: J.B. Lippincott Company, 1982.