

Table 1. Primers used for this study

Gene	Purpose	Sequence (5' to 3')
<i>ACT1</i>	Real-time PCR (F)	TGCACTTCCACATGCTATCC
(At2G37620)	Real-time PCR (R)	GAGCTGGTTTTGGCTGTCTC
<i>Hsp15.7-CI</i>	Real-time PCR (F)	TCAACGGCTCTGATTGATTG
(At5g37670)	Real-time PCR (R)	ACTTCCACCACCGGAAAAAAG
<i>Hsp17.4-CIII</i>	Real-time PCR (F)	CCCGGAATTTCAAATCAGATA
(At1g54050)	Real-time PCR (R)	GCCGTTACAGAAGCCATATCA
<i>Hsp101</i>	Real-time PCR (F)	AATTGAACCTCACGCCTTGG
(At1g74310)	Real-time PCR (R)	CTGCCTCCTGCAAAGAAAAC
<i>Hsp70</i>	Real-time PCR (F)	TAAGGTTCTTCCGGTCCAG
(AT3G12580)	Real-time PCR (R)	CTTGACGCTGAGAGTCGTG
<i>CPK3</i>	pRE8(XVE) (F)	GAGACTCGAGATGGGCCACAGACAGCAAG
(At4g23650)	pRE8(XVE) (R)	GAGAAGTAGTCTAAATCCACGGATGATTAGCAC
<i>CPK13</i>	pRE8(XVE) (F)	GAGACTCGAGATGGGAAACTGTTGCAGATCTCC
(At3g51850)	pRE8(XVE) (R)	GAGAAGTAGTCTAAGCACTTGCTTGCAGTCAGC
<i>HsfB2a</i>	pGreen0229 (F)	GAGATTCGAAATGAATTCGCCGCCGGTT
(At5g62020)	pGreen0229 (R)	GAGAGGATCCATTACAAACTCTCTGATT

(F) and (R) indicate the forward and reverse primers, respectively.

repressors of target gene expression (Czarnecka-Verner et al. 2000; Czarnecka-Verner et al. 2004). This assumption was, in fact, supported by our findings that *hsfB2a* mutants showed significantly higher transcript levels of *Hsp101* and *Hsp15.7-CI* but not other Hsp genes (*Hsp70* and *Hsp17.4-CIII*) in heated seedlings, compared to heat-stressed WT seedlings, (Figure 2B). In contrast, in unheated plants, *hsfB2a* mutants showed slightly lower gene expression for *Hsp101* and *Hsp15.7-CI*, compared to WT (Figure 2A). No phenotypic changes were, however, found for the mutant seedlings, irrespective of heated or not-heated conditions, when compared to WT (data not shown).

To confirm the *in vivo* function of the CPK-HsfB2a cascade, constitutively active forms of CPK and HsfB2a were co-expressed as cofactors for the transient expression of a reporter (GUS) gene under the control of HSE (six inverted repeats of nGAAn units) in *Nicotiana benthamiana* leaves, in *Agrobacterium tumefaciens*-mediated transient expression (agroinfiltration) assays. As shown in Figure 3, the activity of the reporter gene was increased when it was co-expressed with HsfB2a as effector (20-fold). Moreover, the activation by HsfB2a was not additionally enhanced when a constitutively active form of CPK3 or CPK13 protein, which lacks the junction and calmodulin-like domains and thus no longer shows Ca²⁺ dependency, was co-expressed. According to these results, we concluded that the CPK3/13-phosphorylated HsfB2a does not function in the suppression/activation of Hsp expression and enhancement of thermotolerance. As described above, since class B-Hsfs generally interact antagonistically with A-Hsfs by binding (or competing for binding) to the HSE consensus sequence, A-Hsfs should be investigated as possible additional cofactors in further studies. It should also be examined whether a suite of A-Hsfs, of which we have not been tested for CPK substrate targeting (i.e., HsfA1d,

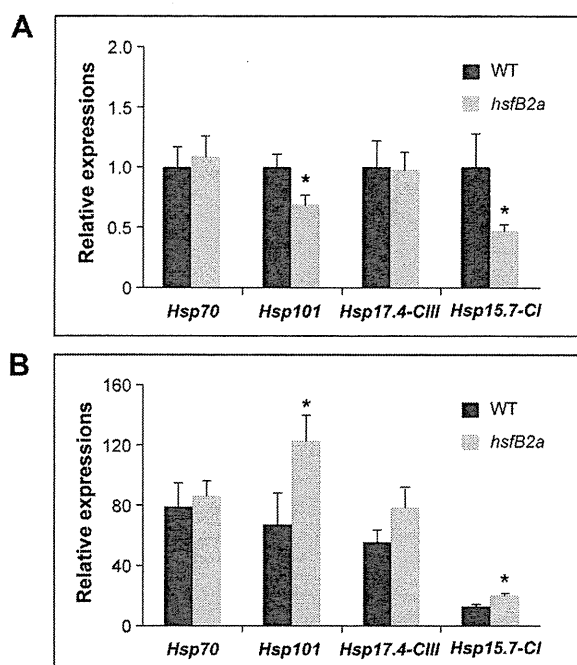


Figure 2. Effects of loss of HsfB2a function on heat response signaling pathway in Arabidopsis seedlings. Transcript levels of Hsp genes in WT and HsfB2a T-DNA insertion mutant (Salk_027578) seedlings before (A) and after (B) heat stress treatment at 40°C for 3 h. Transcript levels of genes were normalized by those of *ACT1* measured in the samples and expressed relative to the normalized transcript levels in the leaves of unheated WT plants. Data represent the mean±SE ($n>5$). An asterisk (*) indicates that the mutant was significantly different from WT for the indicated gene and treatment ($P<0.05$, ANOVA).

HsfA8 (Hsf5), HsfA7a, HsfA7b, HsfA5, HsfA1a (Hsf1) and HsfA3), are directly phosphorylated by CPK3 or CPK13, and then potentially activate the Hsp transcription in an HsfB2a-independent manner.

Our results show that CPK3 and CPK13 function as positive regulators of Hsp gene up-regulation involved in

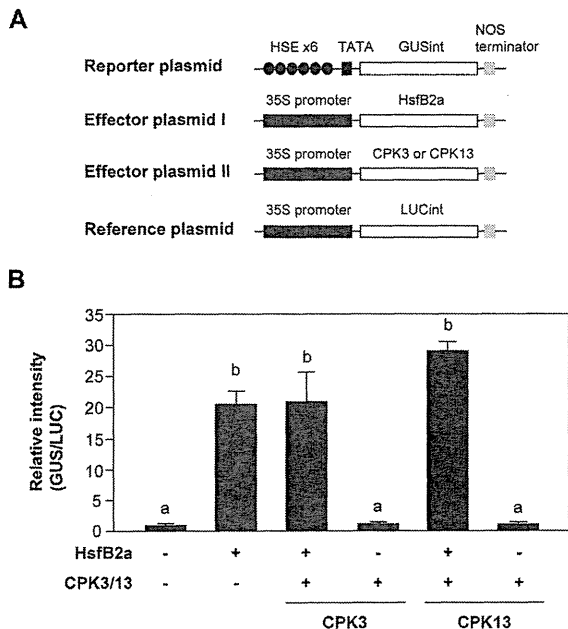


Figure 3. Agroinfiltration assay of a GUS reporter gene under the control of HSE in *Nicotiana benthamiana*. (A) Schematic diagram of the reporter and effector plasmids used in transient assay. An HSE (six inverted repeats of nGAAn units, see underlined sequences) fragment fused to a minimal TATA box: aGAAG cTTCC aGAAC gTTCg aGAAC gTTCg ccc ttc etc tat ata agg aag ttc att tca tt gga gag gac tcc ggt) fragment was fused to a minimal TATA box and a GUS reporter gene including the intron (GUSint). Transient activation of the reporter gene according to co-expressed effector(s), HsfB2a, or truncated variant of CPK3 or CPK13, in *N. benthamiana* leaves was assessed according to a modification of the protocol from (Nagamangala Kanchiswamy et al. 2010). *Agrobacterium*, carrying 35S promoter::LUC including intron (LUCint), was used to normalize for the efficiency of agroinfiltration. (B) Transactivation of a GUS reporter gene under the control of HSE. Data represent the mean \pm SE ($n=5$). Means followed by different small letters are significantly different ($P<0.05$, ANOVA followed by Fisher's PLSD test).

HS tolerance in Arabidopsis. In fact, CPK3 (together with CPK6) is also known to play a role in guard cell ion channel regulation that transduces stomatal abscisic acid signaling (Mori et al. 2006). Moreover, CPK3 has been reported to be associated with the plasma membrane and vacuoles, both depending on its N-terminal myristoylation in the salt stress responses, and thus is able to phosphorylate predominantly membrane-associated proteins (Mehlmer et al. 2010). Since the heat-induced withering of *cpk3* seedlings seem to be due to water-loss (see Figure 1C), a significant portion of CPK3 that function in the heat response signaling pathway may be associated with drought/salt stress-related membrane proteins rather than Hsfs (Arimura and Sawasaki, 2010). Therefore, it is very likely that CPKs contribute to a wide range of central signal transduction responses in plants, whereas their substrate targets are differently involved or function differently.

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References

- Arimura G, Sawasaki T (2010) Arabidopsis CPK3 plays extensive roles in various biological and environmental responses. *Plant Sig Behav* 5: 1263–1265
- Baniwal SK, Bharti K, Chan KY, Fauth M, Ganguli A, Kotak S, Mishra SK, Nover L, Port M, Scharf KD et al. (2004) Heat stress response in plants: a complex game with chaperones and more than twenty heat stress transcription factors. *J Biosci* 29: 471–487
- Czarnecka-Verner E, Pan S, Salem T, Gurley WB (2004) Plant class B HSFs inhibit transcription and exhibit affinity for TFIIB and TBP. *Plant Mol Biol* 56: 57–75
- Czarnecka-Verner E, Yuan CX, Scharf KD, Englich G, Gurley WB (2000) Plants contain a novel multi-member class of heat shock factors without transcriptional activator potential. *Plant Mol Biol* 43: 459–471
- Hu XL, Liu RX, Li YH, Wang W, Tai FJ, Xue RL, Li CH (2010) Heat shock protein 70 regulates the abscisic acid-induced antioxidant response of maize to combined drought and heat stress. *Plant Growth Regul* 60: 225–235
- Klimecka M, Muszyńska G (2007) Structure and functions of plant calcium-dependent protein kinases. *Acta Biochim Pol* 54: 219–233
- Li B, Liu HT, Sun DY, Zhou RG (2004) Ca²⁺ and calmodulin modulate DNA-binding activity of maize heat shock transcription factor in vitro. *Plant Cell Physiol* 45: 627–634
- Li S, Fu Q, Huang W, Yu D (2009) Functional analysis of an Arabidopsis transcription factor WRKY25 in heat stress. *Plant Cell Rep* 28: 683–693
- Liu HT, Gao F, Li GL, Han JL, Liu DL, Sun DY, Zhou RG (2008) The calmodulin-binding protein kinase 3 is part of heat-shock signal transduction in *Arabidopsis thaliana*. *Plant J* 55: 760–773
- Liu HT, Li GL, Chang H, Sun DY, Zhou RG, Li B (2007) Calmodulin-binding protein phosphatase PP7 is involved in thermotolerance in *Arabidopsis*. *Plant Cell Environ* 30: 156–164
- Liu HT, Sun DY, Zhou RG (2005) Ca²⁺ and AtCaM3 are involved in the expression of heat shock protein gene in *Arabidopsis*. *Plant Cell Environ* 28: 1276–1284
- Mehlmer N, Wurzing B, Stael S, Hofmann-Rodrigues D, Csaszar E, Pfister B, Bayer R, Teige M (2010) The Ca²⁺-dependent protein kinase CPK3 is required for MAPK-independent salt-stress acclimation in Arabidopsis. *Plant J* 63: 484–498
- Montero-Barrientos M, Hermosa R, Cardoza RE, Gutiérrez S, Nicolás C, Monte E (2010) Transgenic expression of the *Trichoderma harzianum hsp70* gene increases Arabidopsis resistance to heat and other abiotic stresses. *J Plant Physiol* 167: 659–665
- Mori IC, Murata Y, Yang Y, Munemasa S, Wang YF, Andreoli S,

- Tiriac H, Alonso JM, Harper JF, Ecker JR, et al. (2006) CDPKs CPK6 and CPK3 function in ABA regulation of guard cell S-type anion- and Ca²⁺-permeable channels and stomatal closure. *PLoS Biol* 4: e327
- Nagamangala Kanchiswamy C, Takahashi H, Quadro S, Maffei ME, Bossi S, Berteza C, Atsbaha Zebelo S, Muroi A, Ishihama N, Yoshioka H et al. (2010) Regulation of Arabidopsis defense responses against *Spodoptera littoralis* by CPK-mediated calcium signaling. *BMC Plant Biol* 10: 97
- Sanders D, Pelloux J, Brownlee C, Harper JF (2002) Calcium at the crossroads of signaling. *Plant Cell* 14 Suppl: S401–S417
- Tonsor SJ, Scott C, Boumaza I, Liss TR, Brodsky JL, Vierling E (2008) Heat shock protein 101 effects in *A. thaliana*: genetic variation, fitness and pleiotropy in controlled temperature conditions. *Mol Ecol* 17: 1614–1626
- von Koskull-Döring P, Scharf KD, Nover L (2007) The diversity of plant heat stress transcription factors. *Trends Plant Sci* 12: 452–457
- Wang W, Vinocur B, Shoseyov O, Altman A (2004) Role of plant heat-shock proteins and molecular chaperones in the abiotic stress response. *Trends Plant Sci* 9: 244–252
- Yildiz M, Terzi H (2008) Small heat shock protein responses in leaf tissues of wheat cultivars with different heat susceptibility. *Biologia* 63: 521–525
- Zhang LR, Xing D, Wen F (2009) An on-line multi-parameter analyzing optical biosensor for real-time and non-invasive monitoring of plant stress responses *in vivo*. *Chinese Sci Bull* 54: 4009–4016

Arabidopsis CPK3 plays extensive roles in various biological and environmental responses

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Plant Ca²⁺-dependent protein kinase (CPK) signaling is involved in a wide array of intracellular signaling pathways involved in stomatal movement and plant adaptation to various environmental challenges including drought, salt and cold stress. Arabidopsis CPK3 appears to be extensively involved in such a wide range of aspects, and has been shown to function in mediating the signaling following Ca²⁺ influx after insect herbivory. The results reveal the involvement of CPK3 in the herbivory-induced signaling network through phosphorylating the substrate target HsfB2a (heat shock transcription factor) for transcriptional activation of the plant defensin gene *PDF1.2*. Proteomic studies based on the cell-free protein production system allowed us to mine CPK3 targets more extensively and clarify the nature of multifunctional CPK3.

Plants possess several classes of Ca²⁺-binding sensory proteins, including calmodulins (CaMs), calmodulin-like proteins, calcineurin B-like proteins and CPKs.¹ Especially, CPKs constitute a large family of serine/threonine protein kinases that are broadly distributed in the plant kingdom; for instance, the Arabidopsis genome is predicted to have 34 different CPKs.² Subcellular localization analysis for CPK isoforms from Arabidopsis demonstrated, for example, that AtCPKs 1, 7, 8, 9, 16, 21 and 28 are membrane-associated and other isoforms (AtCPKs 4 and 32) are localized with a nuclear/cytosolic, nuclear and peroxisomal distributions, respectively.^{3,4} Hence, a variety of features make CPKs optimally adapted to control

diverse responses to elevated Ca²⁺, including differences in Ca²⁺ responsiveness, localization and substrate preference.⁵

Arabidopsis CPK3 is of special interest, since this kinase is able to act in a manner that depends on a sizable array of (sub)cellular localizations and potentially specific developmental and environmental conditions (Fig. 1). This CPK is associated with the plasma membrane and vacuoles, both depending on its N-terminal myristoylation in the salt stress response, and thus is able to phosphorylate predominantly membrane-associated proteins.⁶ In guard cells, CPK3 appears to act in the phosphorylation of plasma membrane S-type anion channels for the Ca²⁺-reactive stomatal closure response controlled by abscisic acid (ABA) signaling.⁷ Moreover, our recent research showed that CPK3 also phosphorylates three nuclear/cytosolic transcription factors [jasmonic acid/ethylene-inducible APE/ERF domain transcription factor 1 (ERF1),⁸ heat shock transcription factor HsfB2a, and the wound-inducible CZF1/ZFAR1 transcription factor].^{9,10} The CPK3-derived phosphorylation of HsfB2a promotes *PDF1.2* transcriptional activation in the defense response against a feeding generalist herbivore, *Spodoptera littoralis*. ATL2, a member of a multigene family of highly related RING-H2 zinc finger proteins that function as E3 ubiquitin ligases and a potent regulator of *PDF1.2* transcription,¹¹⁻¹³ is also phosphorylated by CPK3. Since herbivory responses in Arabidopsis is not mediated strikingly by the ABA signaling network, additional branches in the herbivory signal transduction network via

Key words: Arabidopsis, Ca²⁺-dependent protein kinase (CPK), herbivore

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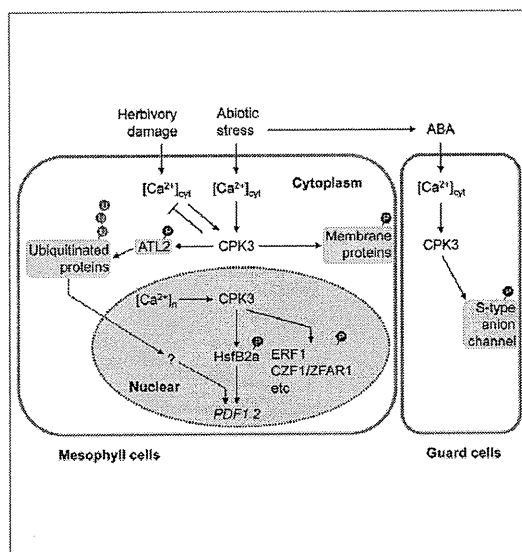


Figure 1. The multifunctional CPK3 signaling required for biotic/abiotic responses.

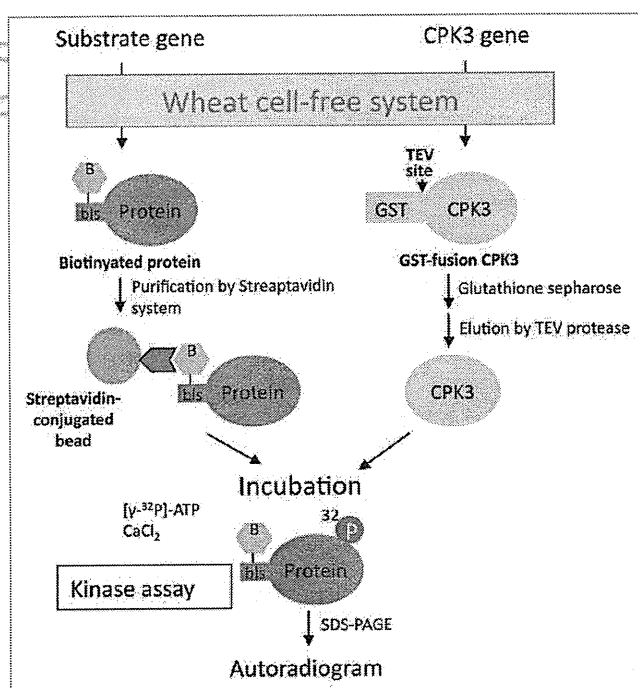


Figure 2. Screening method for the CPK3 substrate identification using the wheat cell-free system.

CPK3 are most likely different from those of the salt stress acclimation as well as the stomatal closure response that are intensively controlled by ABA signaling.^{6,7} In addition, the herbivory-induced cascade is also independent from the jasmonic acid/ethylene-related signaling pathways.¹⁰

Meta Substrate Targeting of CPK3

The identification of protein kinase substrates is important to assess the biological roles of kinases. In addition, recent advances in proteomic technology allow to make use of mass spectrometry MALDI/

TOF for identification of the significant phosphorylation sites of substrates,¹⁴ but this method still leaves us far from understanding the functional nature of kinases. Hence, we still need to rely on a wide array of methods to approach the roles of kinases.

For identification of target substrates of protein kinases, a protein library composed of recombinant proteins is a useful platform. Recent protein production technology using a wheat cell-free system allowed the preparation, for example, of 13,000 human proteins as a protein library.¹⁵ Through the screening of Arabidopsis CPK substrates, four CPK3-phosphorylated targets were successfully mined from 100 nuclear and cytosolic proteins that were conveniently prepared with the cell-free system using 96-well plates (Fig. 2).¹⁰ These results further confirm that this system is a powerful tool to discover the nature of multifunctional protein kinases such as CPK3.

References

- Sanders D, Pelloux J, Brownlee C, Harper JF. Calcium at the crossroads of signaling. *Plant Cell* 2002; 14:401-17.
- Cheng SH, Willmann MR, Chen HC, Sheen J. Calcium signaling through protein kinases. The Arabidopsis calcium-dependent protein kinase gene family. *Plant Physiol* 2002; 129:469-85.
- Dammann C, Ichida A, Hong B, Romanowsky SM, Hrabak EM, Harmon AC, et al. Subcellular targeting of nine calcium-dependent protein kinase isoforms from Arabidopsis. *Plant Physiol* 2003; 132:1840-8.
- Choi HI, Park HJ, Park JH, Kim S, Im MY, Seo HH, et al. Arabidopsis calcium-dependent protein kinase AtCPK32 interacts with ABF4, a transcriptional regulator of abscisic acid-responsive gene expression and modulates its activity. *Plant Physiol* 2005; 139:1750-61.
- Billker O, Lourido S, Sibley LD. Calcium-dependent signaling and kinases in apicomplexan parasites. *Cell Host Microbe* 2009; 5:612-22.
- Mehlmer N, Wurzingner B, Stael S, Hofmann-Rodriguez D, Csaszar E, Pfister B, et al. The Ca²⁺-dependent protein kinase CPK3 is required for MAPK-independent salt-stress acclimation in Arabidopsis. *Plant J* 2010; 63:484-98.
- Mori IC, Murata Y, Yang Y, Munemasa S, Wang YF, Andreoli S, et al. CDPKs CPK6 and CPK3 function in ABA regulation of guard cell S-type anion- and Ca²⁺-permeable channels and stomatal closure. *PLoS Biol* 2006; 4:327.
- Lorenzo O, Solano R. Molecular players regulating the jasmonate signalling network. *Curr Opin Plant Biol* 2005; 8:532-40.
- Cheong YH, Chang HS, Gupta R, Wang X, Zhu T, Luan S. Transcriptional profiling reveals novel interactions between wounding, pathogen, abiotic stress and hormonal responses in Arabidopsis. *Plant Physiol* 2002; 129:661-77.

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10. Nagamangala Kanchiswamy C, Takahashi H, Quadro S, Maffei ME, Bossi S, Berrea C, et al. Regulation of Arabidopsis defense responses against *Spodoptera littoralis* by CPK-mediated calcium signaling. *BMC Plant Biol* 2010; 10:97.
 11. Salinas-Mondragón RE, Garcidueñas-Piña C, Guzmán P. Early elicitor induction in members of a novel multigene family coding for highly related RING-H2 proteins in *Arabidopsis thaliana*. *Plant Mol Biol* 1999; 40:579-90.
 12. Aguilar-Henonin L, Bravo J, Guzmán P. Genetic interactions of a putative *Arabidopsis thaliana* ubiquitin-ligase with components of the *Saccharomyces cerevisiae* ubiquitination machinery. *Curr Genet* 2006; 50:257-68.
 13. Serrano M, Guzmán P. Isolation and gene expression analysis of *Arabidopsis thaliana* mutants with constitutive expression of *ATL2*, an early elicitor-response RING-H2 zinc-finger gene. *Genetics* 2004; 167:919-29.
 14. Sugiyama N, Nakagami H, Mochida K, Daudi A, Tomita M, Shirasu K, et al. Large-scale phosphorylation mapping reveals the extent of tyrosine phosphorylation in Arabidopsis. *Mol Syst Biol* 2008; 4:193.
 15. Goshima N, Kawamura Y, Fukumoto A, Miura A, Honma R, Satoh R, et al. Human protein factory for converting the transcriptome into an in vitro-expressed proteome. *Nat Methods* 2008; 5:1011-7.

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Characterization of a caspase-3-substrate kinome using an N- and C-terminally tagged protein kinase library produced by a cell-free system

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Caspase-3 (CASP3) cleaves many proteins including protein kinases (PKs). Understanding the relationship(s) between CASP3 and its PK substrates is necessary to delineate the apoptosis signaling cascades that are controlled by CASP3 activity. We report herein the characterization of a CASP3-substrate kinome using a simple cell-free system to synthesize a library that contained 304 PKs tagged at their N- and C-termini (N/Ctagged PKs) and a luminescence assay to report CASP3 cleavage events. Forty-three PKs, including 30 newly identified PKs, were found to be CASP3 substrates, and 28 cleavage sites in 23 PKs were determined. Interestingly, 16 out of the 23 PKs have cleavage sites within 60 residues of their N- or C-termini. Furthermore, 29 of the PKs were cleaved in apoptotic cells, including five that were cleaved near their termini *in vitro*. In total, approximately 14% of the PKs tested were CASP3 substrates, suggesting that CASP3 cleavage of PKs may be a signature event in apoptotic-signaling cascades. This proteolytic assay method would identify other protease substrates.

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Subject Category: Immunity

On the basis of the corresponding genetic sequences, >500 human and mouse proteolytic enzymes have been predicted.¹ This number is comparable with that found for protein kinases (PKs), which are the main signal-transduction enzymes.^{2,3} Proteases are involved in the maturation, localization, stabilization, and complex formation of proteins, and in many biological processes, for example, normal development,^{4,5} cancer,^{6,7} infectious diseases,⁸ and cell death.⁹ Therefore, it is important to be able to identify protease substrates using simple assays.

Apoptosis requires the action of many different proteins that participate in apoptotic cell-signaling pathways.¹⁰ Caspases and PKs are critical components of growth and apoptosis signaling pathways.^{2,10} Large-scale analyses of the biological networks involving PKs and caspases are vital for the elucidation of apoptosis signaling pathways. Recent whole-cell proteomic studies that used mass spectrometry attempted to identify substrates of caspases that are involved in apoptosis and have shown that the percentage of PKs found as caspase substrates during apoptosis is 3–6% of ~300.^{11,12} However, cellular protein expression levels may have biased the results.¹³ Furthermore, it is difficult to identify specific pairs of proteases and substrates because numerous cleavage events occur simultaneously in cells. Therefore, an *in vitro* approach that could identify specific proteases and their corresponding substrates would complement cell-based approaches. A diagram, derived from a comprehensive *in vitro* study, that illustrates the relationships between

caspases and their PK substrates would help clarify the signal-transduction events that occur during apoptosis.

A collection of recombinant proteins, that is, a protein library, is needed to screen a large number of protein substrates. In addition, to screen a protein library comprehensively two *in vitro* high-throughput methods – one for protein synthesis and one for the detection of the targeted biochemical reaction – are required. Recently, we developed an automated protein synthesis system that uses a wheat cell-free system.^{14–16} Using this system, we were able to synthesize many human and Arabidopsis PKs.^{17,18} Recent work by others suggested that the wheat cell-free system could produce 13364 human proteins, which, because of the large number of proteins involved, represents an *in vitro*-expressed proteome.¹⁹ We also recently developed a method to label monobiotin proteins that had been synthesized in the wheat cell-free system.²⁰ These monobiotin-labeled proteins were then used directly – without purification – to detect protein ubiquitination²¹ and an autoantibody in the serum.²² As the procedures used with many commercially available detection kits depend on biotin–streptavidin interactions, our purification-free, synthesis/biotin-labeling method provides a simple and highly specific system that can be used for biochemical analyses.

Caspase-3 (CASP3) cleaves many different proteins,^{23,24} and its action *in vivo* irreversibly induces apoptosis. For the study reported herein, we delineated a CASP3-substrate kinome using a simple luminescent-based detection method

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Keywords: caspase; protein kinases; apoptosis; cell-free protein synthesis; protein library

Abbreviations: CASP3, caspase 3; PK, protein kinase; N/Ctagged, N- and C-terminally tagged; TD, terminal detection

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to screen an N- and C-terminally tagged (Ntagged) PK library produced in the wheat cell-free system. This comprehensive characterization of a CASP3-substrate kinome is a resource that can be used to understand the roles of PKs in apoptosis.

Results

Generation of an Ntagged PK library used to identify CASP3 PK substrates. To identify PKs that are substrates of CASP3, we first made a library consisting of 248 human and 56 mouse PKs (Supplementary Table S1). The nucleotide sequences for the Flag-tag and the biotin ligation site (bls) were added upstream and downstream, respectively, of the PK open-reading frame by PCR incorporation of Gateway recombination tags. Each PCR product (attB1-Flag-PK-blb-attB2) was inserted into a pDONR221 vector using the Gateway BP Clonase II system (upper panel, Figure 1). The Flag-PK-blb nucleotide sequences from the *Escherichia coli* cultures were used without purification to construct, by split-primer PCR, the DNA templates for protein synthesis.¹⁴ The Ntagged PK library (304 PKs) was produced using an automated protein synthesizer (GenDecoder 1000; CellFree Sciences Co., Ltd., Matsuyama, Japan), with biotin and biotin ligase added into the synthesis mixtures for monobiotin labeling at the bls.^{20,21} That the members of the protein library were Ntagged was confirmed by immunoblotting with anti-Flag antibodies and Alexa488-labeled streptavidin.

To assess the suitability of the designed PKs to act as CASP3 substrates, we used Ntagged p21-activated kinase 2 (PAK2), which is a known CASP3 substrate,²⁵ as the test case. The biotinylated Ntagged-PAK2 (Flag-PAK2-blb~biotin) was treated with CASP3 and cleavage of PAK2 was confirmed by immunoblotting with Alexa488-conjugated streptavidin (Figure 2a). In addition, the cleavage site (₃₁₉DELD↓_{S323}), determined by amino-acid sequencing, was found to be the same as that reported previously.²⁵ (The arrow indicates the hydrolytic bond.)

A luminescent assay to detect PK substrates of CASP3.

A schematic of the assay used to monitor cleavage of the Ntagged PKs by CASP3 is shown in Figure 1. The PK construct is first incubated with CASP3. If the construct contains a sequence that can be cleaved by CASP3, cleavage occurs. Acceptor and donor beads are then added. The Flag-tag binds a protein A-conjugated acceptor bead via an anti-Flag antibody, and the biotin bound to the C-terminus of the PK construct binds a streptavidin-conjugated donor bead. If an acceptor bead is in close contact with the donor bead, as is the case when the construct is not a CASP3 substrate and both beads are therefore bound intramolecularly, the system luminesces. However, if CASP3 had cleaved the Ntagged PK, luminescence is suppressed because the beads are no longer in close contact. As a proof-of-concept experiment, cleavage of the test PK, Ntagged PAK2, was assessed

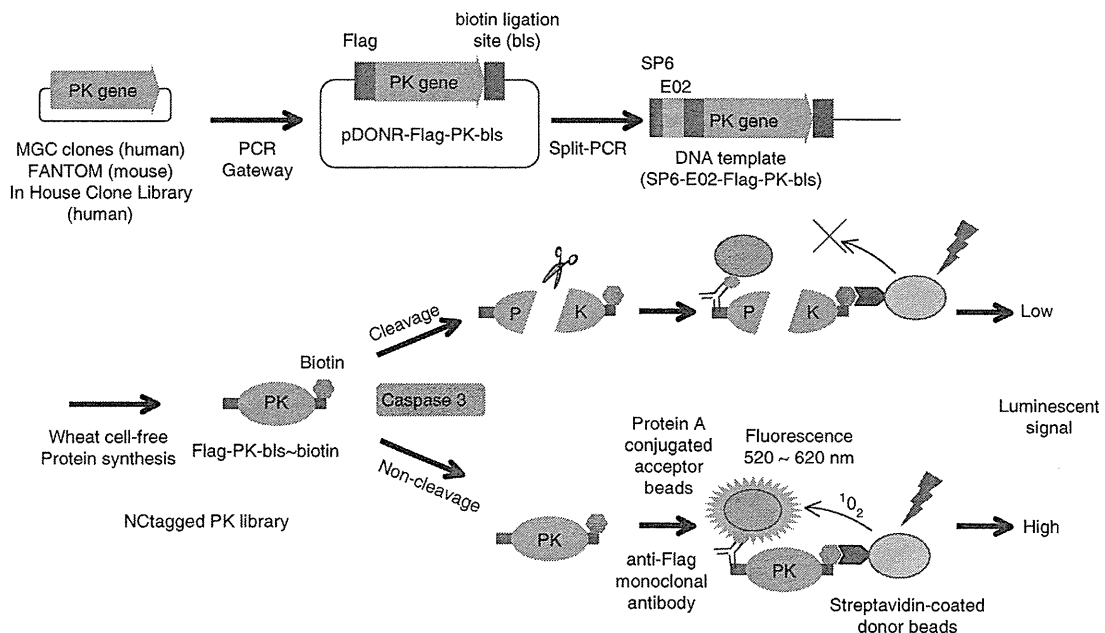


Figure 1 Schematics of the DNA template construction and the CASP3-substrate-screening assay. *Protein kinase (PK)* genes were obtained from the human MGC and mouse FANTOM libraries, and from a library of *PK* genes that we had cloned. The *PK* genes were PCR amplified with the Flag and the biotin ligation site (bls) tags added to the upstream and downstream ends, respectively. The modified genes were each inserted into a Gateway pDONR221 vector (pDONR-Flag-PK-blb) and DNA templates (SP6-E02-Flag-PK-blb) were constructed by split-primer PCR and then expressed in the wheat cell-free protein synthesis system that included biotin ligase and D-biotin to give Flag-PK-blb~biotin constructs. The Flag and biotin tags were bound to protein A-conjugated acceptor beads via an anti-Flag antibody and streptavidin-conjugated donor beads, respectively. An intact complex luminesced strongly, whereas after CASP3 cleavage and dissociation of the protein fragments, the luminescence was abolished or reduced

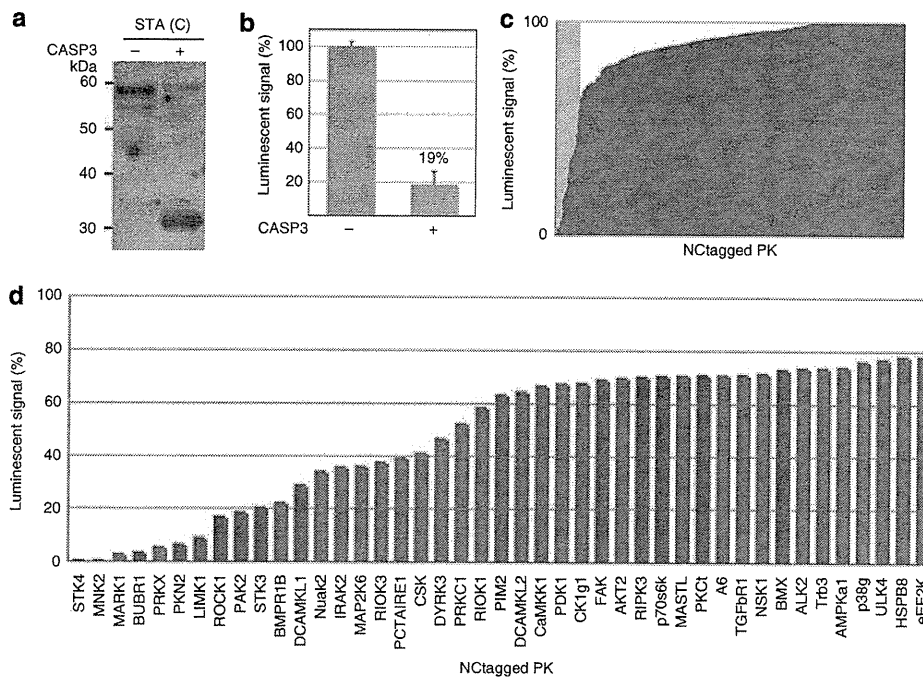


Figure 2 Screening of CASP3-cleaved PK substrates from the NTagged PK library. (a) Immunoblot of NTagged PAK2 that had been incubated in the presence (+) or absence (-) of CASP3. Alexa488-labeled streptavidin (STA(C)) was used for detection. (b) Detection of CASP3-cleaved NTagged PAK2 using the AlphaScreen system. The luminescence for the control (no CASP3) was set to 100%. The value of 19% indicated that most of the NTagged PAK2 was cleaved by CASP3. Each value is the mean of three independent experiments, and the uncertainty is reported as the standard deviation. (c) Luminescent signals remaining after *in vitro* CASP3 treatment of NTagged PKs that had been synthesized in the wheat cell-free system. The x axis lists the NTagged PKs in ascending order of their luminescent signals after CASP3 treatment. (d) NTagged PKs that returned luminescent signals of <78% of the control values. The plot contains the data of (c) within the green rectangle. Red bars are for PKs that were known to be substrates of CASP3 before this report

using this system. CASP3 treatment decreased the luminescent signal to $19 \pm 7\%$ that of the control (no CASP3; Figure 2b). Therefore, the system could detect CASP3 cleavage and can replace conventional immunoblotting procedures.

Screening of the CASP3-substrate kinome. Using the luminescent system, 304 NTagged PKs were screened. The level of luminescence after CASP3 treatment is reported as the percentage of the corresponding control (no CASP3; Figure 2c and d). Thirteen of the NTagged PKs for which luminescence was low after CASP3 treatment are known CASP3 substrates.^{23,24,26} The smallest and largest luminescent values were for STK4 (1%) and BMX (73%), respectively; we therefore examined the physical states of the PKs that had been treated with CASP3 and had associated luminescence values of $\sim 80\%$ by immunoblotting with anti-Flag antibodies and Alexa488-streptavidin to detect the N- and C-termini of the NTagged PKs, respectively. This 'terminal detection' (TD) immunoblot assay identified 43 NTagged PKs that had been cleaved (Supplementary Table S1). In addition to the 13 PKs that were known to be CASP3 substrates, 30 previously unidentified PK that were substrates of CASP3 were found (Figure 3 and Table 1). In addition, because the apparent molecular weights of the N- and C-terminal fragments could be estimated from their positions in the TD immunoblot, the

CASP3 cleavage sites could be predicted (red arrowheads, Figure 3). For MASTL, the signal on the immunoblot with Alexa488-conjugated streptavidin was not detectable, probably indicating that the efficiency of biotinylation in MASTL proteins might be too low to detect for the immunoblot. Luminescent signal of this clone was also very low (see Supplementary Table S1).

A comparison of the luminescent and immunoblot data correlated a luminescent signal of <78% with a positive immunoblot result. Forty-eight PK constructs with luminescent signals >78% were tested and returned negative immunoblot results (Supplementary Table S1). Therefore, a luminescent signal of $\sim 78\%$ is the apparent divisor between PKs that can be cleaved by CASP3 and those that cannot be cleaved.

***In vivo* identification of the PKs that were identified as CASP3 substrates by the luminescent assay.** We investigated whether the newly identified PKs that were substrates of CASP3 were cleaved in HeLa cells that had been induced to undergo apoptosis by TNF α plus cycloheximide (TNF α)²⁷ or anti-Fas antibody (anti-Fas).²⁸ The genes encoding these PKs were each inserted into the transfection vector, pDEST26, using the Gateway system and expressed as (His)₆-PK-Flag constructs. We were able to detect all expressed PK constructs, except DYRK3, by immunoblotting with anti-(His)₆ or anti-Flag antibodies.

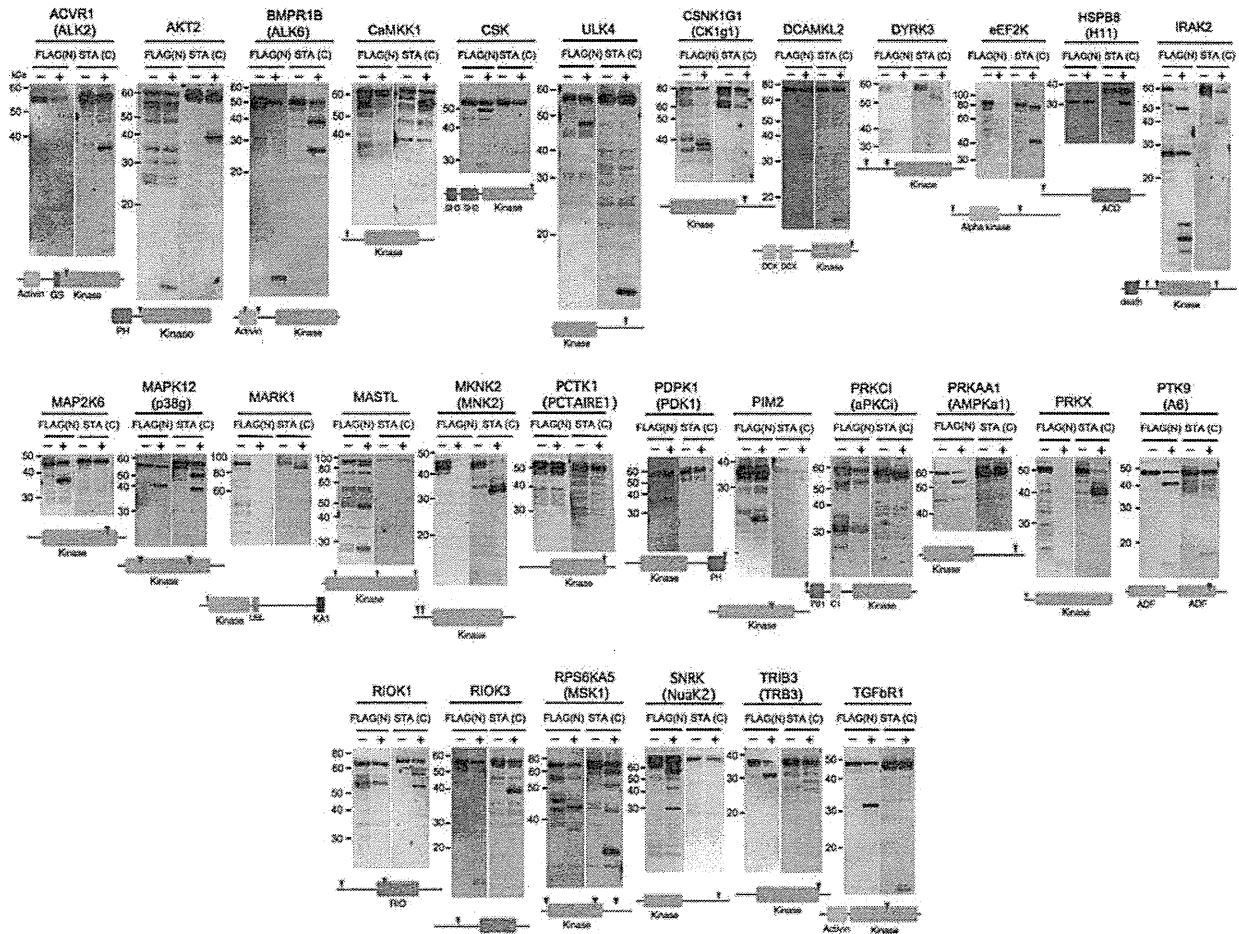


Figure 3 *In vitro* cleavage of NCTagged PKs by CASP3. The NCTagged PKs that had been incubated in the presence (+) or absence (-) of CASP3 and their cleavage products were detected using anti-Flag antibodies (FLAG(N)) and Alexa488-conjugated streptavidin (STA(C)), which bound to the N- and C-termini of the PK constructs, respectively. The cartoons of the proteins that are under the lanes show the locations of the conserved domains (colored boxes) and the predicted cleavage sites (red arrowheads). The conserved domains that are found in the Conserved Domains Database (<http://www.ncbi.nlm.nih.gov/cdd>) are: ACD, alpha-crystallin domain; activin, conserved domain for activin members; ADF, actin depolymerization factor/cofilin-like domain; α -kinase, conserved kinase domain for the α -kinase family; C1, phorbol esters/diacylglycerol binding domain; DCX, doublecortin domain; death, death domain; GS, GS motif; KA1, kinase-associated domain; kinase, catalytic domain of protein kinase; PB1, Phox and Bem1p domain; PH, pleckstrin homology domain; RIO, catalytic domain of eukaryotic RIO kinase family; SH2, src homology 2 domain; SH3, src homology 3 domain; UBL, ubiquitin-like domain

Notably, they were detected as cleavage products and/or were found in smaller amounts when the cells had been induced to undergo apoptosis than when apoptosis had been inhibited by z-VAD-FMK (Figure 4a and b). Furthermore, apoptosis-induced cleavage of four endogenous PKs was found by immunoblotting with commercially available antibodies against the endogenous PKs (Figure 4c). These *in vivo* experiments validated the underlying concept of our *in vitro* cell-free system as the *in vivo* system found all of the PKs identified by the *in vitro* system.

Characterization of the CASP3 cleavage sites in the newly identified PK substrates. We characterized the CASP3 cleavage sites in the newly identified PK substrates. As the positions of the cleaved PK fragments in the TD immunoblot could be used to estimate the size of the cleaved fragments and because the antibodies could be

used to identify whether the fragments were derived from the N- or C-terminal regions of the PKs, we could predict the approximate positions of the CASP3 cleavage sites (red arrowheads, Figure 3). Each NCTagged PK that was a substrate for CASP3 was synthesized in the cell-free system and purified using Streptavidin Magnesphere Paramagnetic beads. Their C-terminal fragments that bound to the beads were recovered after CASP3 cleavage and their sequences were determined. Using this approach, the cleavage sites of ACVR1, AKT2, BMPR1B, CaMKK1, HSPB8, MAPK12, MKNK2(D58), PDPK1, PRKCI, PRKX, RIOK1, RIOK3, and RPS6KA5 were determined. We then attempted to determine the cleavage sites of the remaining PKs by other methods.

The NCTagged PKs that had low biotin-labeling efficiencies and were cleaved near their C-termini were genetically modified by the addition of a glutathione-S-transferase (GST) fragment at their C-termini to facilitate recovery with

Table 1 Characteristics of the newly identified CASP3 PK substrates

Symbols	Kinome names	Groups	Clone origin	AA ^a	Cleavage sequence	Cleavage sites	Methods ^b	Conservation ^c	Smallest frag. ^d	<i>In vivo</i> cleavages ^e
ACVR1	ALK2	TKL	Hs	509	IASD↓M	269	NT	Yes	C240	Yes
AKT2	AKT2	AGC	Mm	481	DAMD↓Y	121	NT	Yes	N121	Yes
BMPR1B	ALK6	TKL	Hs	502	CSTD↓G	50	NT	Yes	N50	Yes
					DFVD↓G	120	NT	Yes		
CaMKK1	CaMKK1	Other	Hs	520	EEAD↓G	32	NT	Yes	N32	Yes
CSK	CSK	TK	Mm	450	DAPD↓G	409	MS	Yes	C41	Yes
CSNK1G1	CK1g1	CK1	Mm	459	VHVD↓S	343	MU	Yes	C116	Yes
eEF2K	eEF2K	Atypical	Hs	725	EGVD↓G	14	MU	Yes	N14	Yes
					DHLD↓N	430	MU	Yes		
HSPB8	H11	Atypical	Hs	196	MAD↓G	3	NT	Yes	N3	Yes
MAP2K6	MAP2K6	STE	Mm	334	DFVD↓F	289	MU	Yes	C45	Yes
MAPK12	p38g	CMGC	Hs	367	SAVD↓G	46	NT	Yes	N46	Yes
MARK1	MARK1	CAMK	Hs	795	SATD↓E	52	MU	Yes	N52	Yes
MKNK2	MNK2	CAMK	Hs	414	DQPD↓H	32	MU	No	N32	Yes
					DIPD↓A	58	NT	Yes		
PDPK1	PDK1	AGC	Hs	556	SHPD↓A	552	NT	Yes	C4	Yes
PIM2	PIM2	CAMK	Hs	334	TDFD↓G	198	MU	Yes	C113	Yes
PRKAA1	AMPKa1	CAMK	Hs	550	TSLD↓S	520	MS	Yes	C30	Yes
PRKCI	aPKCi	PKC	Hs	596	TQRD↓S	6	NT	Yes	N6	Yes
PRKX	PRKX	AGC	Hs	358	ETPD↓G	25	NT	No	N25	Yes
RIOK1	RIOK1	Atypical	Mm	568	EKDD↓I	37	NT	Yes	N37	Yes
RIOK3	RIOK3	Atypical	Hs	516	DTRD↓D	139	NT	Yes	N139	Yes
RPS6KA5	MSK1	AGC	Hs	549	DGGD↓G	20	NT	Yes	N20	Yes
					DELD↓V	344	NT	Yes		
					TEMD↓P	356	NT	Yes		
SNARK	NuaK2	CAMK	Hs	628	VSED↓S	546	MU	Yes	C82	Yes
TRIB3	TRB3	CAMK	Hs	358	VVPD↓G	338	NT	Yes	C20	Yes
ULK4	ULK4	Other	Hs	580	SQID↓S	473	MU	Yes	C107	Yes
DCAMKL2	DCAMKL2	CAMK	Hs	695	—	—	—	—	—	Yes
DYRK3	DYRK3	CMGC	Hs	568	—	—	—	—	—	ND
IRAK2	IRAK2	TKL	Mm	622	—	—	—	—	—	Yes
MASTL	MASTL	AGC	Hs	879	—	—	—	—	—	Yes
PCTK1	PCTAIRE1	CMGC	Hs	496	—	—	—	—	—	Yes
PTK9	A6	Atypical	Hs	384	—	—	—	—	—	Yes
TGFBR1	TGFBR1	TKL	Hs	426	—	—	—	—	—	Yes

Abbreviations: Hs, human clone; Mm, mouse clone; MS, mass spectroscopy; MU, mutation; ND, not determined; NT, N-terminal sequencing. ^aLength of amino acids. ^bMethods for determination of cleavage site. ^cVery similar site conserving the Asp (D) of the hydrolytic bond was found between human and mouse PKs (Yes), whereas no similar sites was done (No). ^dThe smallest N (N) or C (C) fragment in the cleaved PKs. Number is the length of amino acids of the fragment. ^eData from Figure 4.

glutathione Sepharose 4B beads after CASP3 cleavage. CASP3 cleavage of the PK-GSTs produced the same size N-terminal fragments as those of the corresponding CASP3-cleaved NTagged PKs, indicating that the GST tags did not alter the positions of the cleavage sites. In addition, the sequences of the cleaved c-src tyrosine kinase (CSK) and AMP-activated kinase-α1 (AMPKa1) fragments were determined using MALDI/TOF-MS. Other PK constructs that were synthesized in small amounts were subjected to D→A mutagenesis to determine their cleavage sites. In total, 28 cleavage sites in 23 PKs were identified (Table 1). Identical or similar cleavage sites were found in the corresponding human and mouse PKs, except for those of PRKX (Supplementary Table S2). (Sequence analysis showed that mouse PRKX does not have the N-terminal region that is found in human PRKX.) Therefore, the CASP3-substrate kinome may be highly conserved in mammals.

We also analyzed the common sequence attributes among the 28 cleavage sites and found that CASP3 prefers the sequence, DXXD↓G (Figure 5a). The consensus PK cleavage site for CASP3 in the MEROPS database is DXXD↓X. In the NTagged PK library, 208 of the 304 PKs contain a DXXDX sequence. However, only 33 PKs were

cleaved by CASP3; therefore, to be cleaved by CASP3, the DXXDX sequence and a structural element –probably accessibility – are required.

Characterization of the newly identified PKs that were cleaved near their N- or C-termini. Interestingly, 16 out of the 23 PKs, for which cleavage sites were characterized, have cleavage sites within 60 residues of their N- or C-termini. We investigated whether these sites were also cleaved *in vivo* when apoptosis was induced by TNFα. For these experiments, CaMKK1, eEF2K, MNK2, AMPKa1, and TRIB3, which were cleaved *in vitro* at D32, D14, D32/D58, D520 (30 residues away from the C-terminus), and D338 (20 residues away from the C-terminus), respectively, were used (Figure 5b and Table 1). Their genes (wild type, WT) were each reconstructed with a V5 tag added at the end opposite the cleavage site. The genes for their D→A mutants (DA), and for the sequences of their longer CASP3-cleaved fragments (C3M), were also constructed and all were expressed in control and in apoptotic cells (Figure 5c). Cleavage of the WT PKs produced long fragments corresponding to C3M in apoptotic cells, whereas z-VAD-FMK blocked cleavage. These cleavages near the N- and

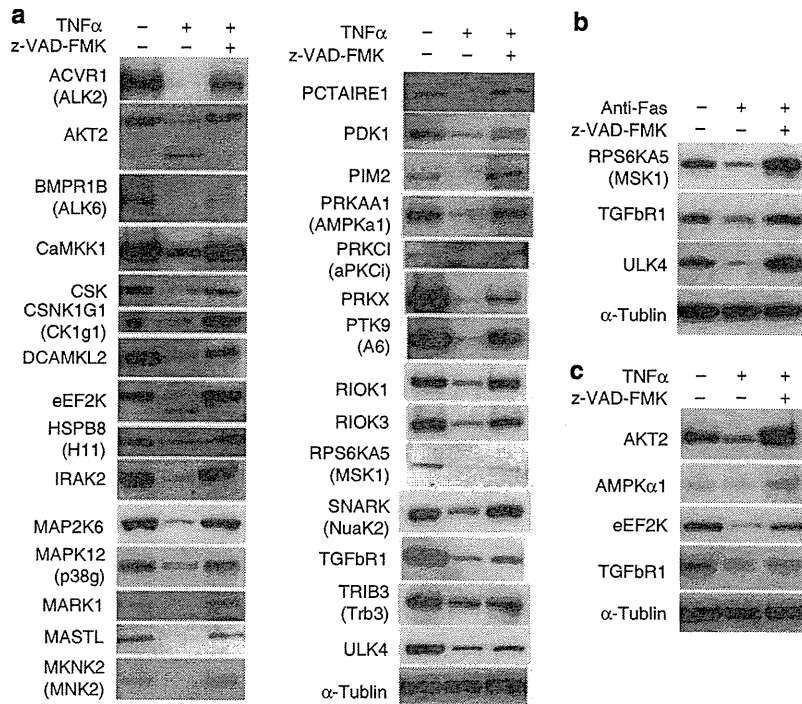


Figure 4 *In vivo* caspase cleavage of the newly identified PK substrates and of endogenous PK substrates. (a) *In vivo* cleavage of the (His)₆-PK-Flag constructs expressed in apoptotic HeLa cells. The cells were treated with DMSO (control) or with TNF α and cycloheximide (TNF α) in the presence and absence of z-VAD-FMK (a CASP3 inhibitor) for 6 h and then lysed. The cell extracts were immunoblotted and the PK constructs were detected with anti-Flag antibodies, except for HSPB8 and MAP2K6. Anti-His tag antibody was used for the two PKs. (b) The cells were transfected with a plasmid of (His)₆-PK-Flag constructs, and treated with DMSO (control), or with anti-Fas antibody (anti-Fas) in the presence and absence of z-VAD-FMK for 6 h and then lysed. Immunoblotting was carried out as (a). (c) HeLa cells were treated as in (a), but were not transfected with a (His)₆-PK-Flag gene. Each endogenous PK was detected using an antibody specific for it. α -Tubulin was used as an internal marker

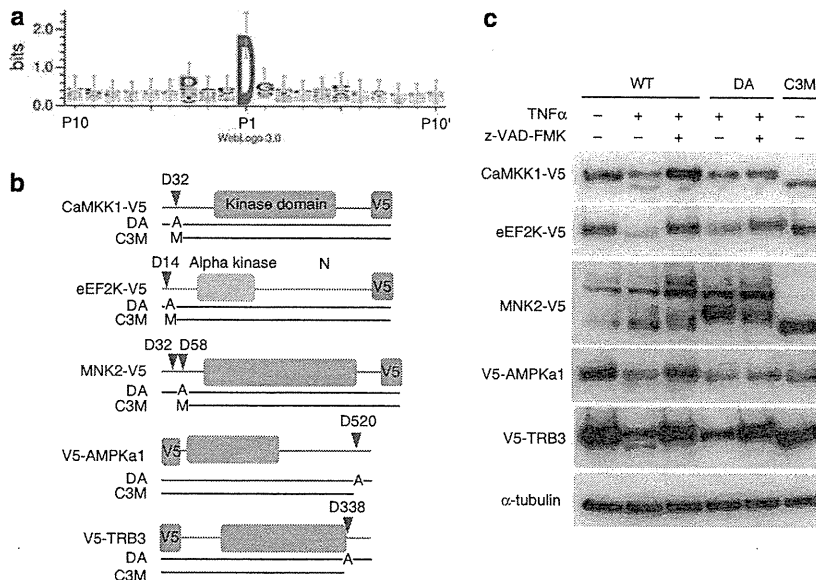


Figure 5 The cleavage site logo and *in vivo* cleavage of five PKs that are cleaved by CASP3 near their N- or C-termini. (a) The 20 residues surrounding the D of the hydrolytic bond in 28 PKs were analyzed using WebLogo, version 3.0.³⁶ (b) Cartoons of five PK sequences that have cleavage sites near their N- or C-termini. The corresponding PKs were used for the experiment shown in (c). The positions of the alanines in the D \rightarrow A mutants (DA) are shown, as are the long fragments (C3M) produced by CASP3 cleavage. V5 tags were fused at the ends farther away from the cleavage sites. The first M in C3M of CaMKK1, eEF2K, and MNK2 indicates a methionine as a start amino acid. (c) Immunoblots of PK-V5s and V5-PKs that had been expressed in apoptotic HeLa cells. The cells were treated with DMSO (control) or with TNF α and cycloheximide in the presence or absence of z-VAD-FMK for 6 h and then lysed. The proteins were blotted and then detected with anti-V5 antibodies. WT indicates a wild-type protein

C-termini of the PKs suggest that CASP3 cleavage may regulate the activity level and/or cellular localization of the PKs, rather than simply inactivate the kinases.

Discussion

In 1995, PITSLRE,²⁹ PKC δ ,³⁰ and DNA-PKcs³¹ were reported as the first PK-type substrates of CASP3. During the next 15 years, 36 additional PKs that can be cleaved by CASP3 were found.^{23,24,26} Notably, these authors showed that CASP3-cleaved PKs abrogate survival signals and accelerate apoptosis. In this study, we identified an additional 30 PKs that can be cleaved by CASP3. In addition, many of the cleavage sites were found in regulatory elements or in the regions near the N- and C-termini, rather than the kinase domain itself. Some of the newly identified CASP3-substrate PKs may be involved in apoptotic signal cascades. Sixteen PKs were shown to be cleaved *in vitro* near their N- or C-termini and at least five of them were also cleaved near their N- or C-termini in apoptotic cells (Figure 5). Using standard immunoblotting, proteins that are cleaved into a large and a small fragment may be overlooked because the mobilities of the large fragment and the intact protein will be nearly identical. Most of the PKs that had been previously reported to be cleaved by CASP3 were identified because the cleaved fragments had very different molecular weights than did the intact PK and were therefore easily detected by SDS-PAGE. Consequently, cleavages near the termini may have been overlooked. Taken together, our results suggest that CASP3 cleavage of some of the members of the CASP3-substrate kinome alters the function of the PKs and thereby signals apoptosis.

For the study reported herein, 304 out of 518 known PKs, synthesized as NTagged PKs, were subjected to the *in vitro* cleavage assay (Supplementary Table S1). The relative number of PKs that were cleaved was ~14%. A total of 69 PKs that are CASP3 substrates are now known, which suggests that at least ~13% of the PKs in the human kinome are targets of CASP3. As ~200 PKs have yet to be tested as CASP3 substrates, an additional 26 PKs (13% of the 200) may be CASP3 substrates. The human genome contains 518 annotated PKs, which have been divided into 10 groups on the basis of their sequence homologies.³ Interestingly, the groups differ in terms of their susceptibilities to CASP3 cleavage (Table 2). Approximately 30% of the PKs in the AGC group are known CASP3 substrates, for example, AKT2, S6K, MSK, PKC, and PDK1. Many of the AGC-type PKs are commonly found in mammalian tissues,³² and their cleavage sites are located in their regulatory domains (Figure 4 and Table 1). Therefore, these abundant PKs may be activated when CASP3 cleaves them and then act as intracellular apoptosis signals. Conversely, CASP3 cleaved only a relatively small number (6~8%) of the PKs in the CMGC group, which includes the kinases of the CDK and CDKL families, and the tyrosine kinase groups. Therefore, most members of these groups may only act indirectly as apoptosis signals after CASP3 activation.

Such ROCK1 and MST1, certain caspase cleavage products, work as apoptosis signaling.^{23,24} In this study, we found at least six new CASP3 cleavage products, derived from AKT2, CaMKK1, eEF2K, MARK1, MNK2, and TRB3,

Table 2 Characteristics of the protein kinases used in this study

Groups	Total ^a	Tested clones	Cleaved clones (new ^b)	Cleaved/test clones (%)
AGC	63	33	10 (6)	30
CAMK	74	52	8 (7)	15
CK1	12	8	1 (1)	13
CMGC	61	39	3 (3)	8
Other	83	46	3 (2)	7
STE	47	25	4 (1)	16
TK	90	51	3 (1)	6
TKL	43	27	6 (4)	22
RGC	5	2	0	0
Atypicals	40	21	5 (5)	24
Total	518	304	43 (30)	14

^aEach number is corresponding to human kinome. ^bNewly PK numbers found in this study.

after 6 h from apoptosis induction (Figures 4 and 5). These cleavage products retain kinase domain, as in the case of ROCK1 and MST1. On the other hand, we could not detect any cleavage products from the other kinases *in vivo*. The reasons are not yet understood. However, recent proteomics approach has shown that the cleaved proteins displayed transient fragments in the apoptotic cells.¹² Further analysis at multiple time points during the apoptotic cascade would be required for detection of the cleavage products from the remaining PKs.

For TRIB3, full-length TRIB3 (D338A) mutant was decreased in apoptotic condition (compared TNF α lane with TNF α plus z-VAD-FMK lane in Figure 5c). However, the mutant could not produce the CASP3 cleavage product found as the shorter form in TNF α lane of WT, indicating that the mutant was not cleaved by CASP3. The mutant was also not cleaved by CASP3 *in vitro* (data not shown). As TRIB3 has been known to receive proteasomal degradation,³³ this unexpected reduction of the mutant TRIB3 in the apoptotic cells may be the effects of cycloheximide and/or caspase-inhibitor treatment on TRIB3 degradation.

Proteases often modify the activities of their targeted protein substrates. Identification of the specific substrate that is cleaved by a protease is necessary if the functions of both the protease and its substrate are to be understood. Proteomic studies have used mass spectrometry to exhaustively identify cellular proteins that have been cleaved by proteases.^{11,12} However, it has been difficult to correlate specific proteases with their substrates because many proteases act at the same time *in vivo*.

Many full-length cDNAs derived from the genes of higher eukaryotes are available from many different sources. These cDNAs are potentially a great DNA template resource for *in vitro* syntheses of proteins. As a protein production system and for the functional analysis of proteins, the wheat cell-free system has many advantages: It can effectively use PCR-generated DNA templates.¹⁴ It is easily adapted to an automated system.¹⁵ It can be used to incorporate a single label into target proteins.²⁰ Its synthesized proteins do not require purification before being assayed, and it has no detectable proteasome activity.²¹ In addition, the screening cost is very low (~US\$1/assay), which for our study translated to 10 cents to produce each NTagged protein and

20 cents for the beads, CASP3, and disposable hardware used in one assay.

In summary, we showed that an NTagged PK library synthesized in a cell-free system could be used to characterize a CASP3-substrate kinome. Analysis of the CASP3-cleavage sites indicated that CASP3 cleavage of PKs depends on both primary and tertiary structure. Almost all of the PK substrates that we identified *in vitro* were also identified *in vivo*. Systems similar to that used herein could be used to screen other protease substrates.

Materials and Methods

General. The following procedures have been described:^{14–16,20–22,34} wheat cell-free protein production; split-primer PCR synthesis of the DNA templates; parallel syntheses of mRNAs and their translated proteins; and measurements of the amounts of protein synthesized using densitometer scans of Coomassie brilliant blue-stained proteins or of radiolabeled proteins.

Construction of DNA templates for the expression of a PK protein library. The cloned genes encoding the PKs used in this study are listed in Supplementary Table S1. Their open-reading frames (without stop codons) were modified in two steps using PCR and the primers S1 (5'-CCACCCACCACACCAatg(n)₁₆-3') and T1 (5'-TCCAGCACTAGCTCCAGA(n)₁₉-3') (lowercase letters indicate nucleotides of the gene) for the first step, and the primers attB1-Flag-S1 (5'-GGGGACAAGTTTGTACAAAAAAGCAGGCTTCATGGACTACAAGGATGACGATGACAAGCTCCACCCACCACCACCAATG-3') and T1-bis-STOP-attB2-anti (5'-GGGGACCACCTTTGTACAAGAAAGCTGGGTTTATTCGTGCCACTC GATCTTCTGGGCCTCGAAGATGCTTTCAGCCGCTCCAGCACTAGCTCCA GA-3') for the second step. The PCR-modified genes were each inserted into a pDONR221 vector using the Gateway BP Clonase II enzyme mix (Invitrogen, Carlsbad, CA, USA) to give pDONR-Flag-PK-bis vectors. *Escherichia coli* cells were transformed with these vectors and then cultured in wells of a 96-well plate that contained GYT medium (10% (v/v) glycerol, 0.125% (w/v) yeast extract, and 0.25% (w/v) tryptone) for 48 h without shaking. DNA templates for mRNA and protein expression were constructed using split-primer PCR¹⁴ in two steps. For the first step, the pDONR221-Flag-PK-bis plasmids that had not been isolated from the *E. coli* cells, and the primers pDONR221-1st_4080 (5'-ATCTTTTCTACGGGTCTGA-3') and deSP6E02-Flag-S1 (5'-GGTGACACTATAGAACTCACCTATCTCTACACAAAACATTTCCCTACATACAACCTTCAACTTCTATTATGGACTACAA GGATGACGATGACAAGCTCCACCCACCACCACCAATG-3') were used, and for the second step, the amplified sequences of the first step and the primers SPu (5'-GGGTAGCATTAGGTGACACT-3') and pDONR221-2nd_4035 (5'-ACGTTAA GGGATTTGGTCA-3') were used to give SP6-E02-Flag-PK-bis DNA templates. (The E02 sequence is a translational enhancer,³⁵ and the SP6 sequence is an SP6 RNA polymerase promoter.)

Cell-free protein synthesis. Cell-free protein synthesis used the reagents of an ENDEXT Wheat Germ Expression S Kit according to the manufacturer's instructions (CellFree Sciences Co., Ltd.), the bilayer translation method,^{15,16,34} and a robotic synthesizer (GenDecoder 1000; CellFree Sciences). Each DNA template was transcribed by SP6 RNA polymerase, then precipitated with ethanol, and collected by centrifugation (15 000 r.p.m. for 5 min., R10H rotor; Hitachi). Each mRNA (~30–35 μg) was washed with 75% ethanol, added into a translation mixture, and translated in the bilayer mode³¹ with the following modifications. The translation mixture (25 μl) (bottom layer) contained 60 A₂₆₀/ml of ENDEXT wheat germ extract, 1 × SUB-AMIX (24 mM Hepes-KOH, pH 7.8, 1.2 mM ATP, 0.25 mM GTP, 16 mM creatine phosphate, 2.5 mM DTT, 0.4 mM spermidine, 0.3 mM each of the 20 amino acids, 2.8 mM magnesium acetate, 100 mM potassium acetate), 2 μg creatine kinase (Roche Applied Science, Indianapolis, IN, USA), 500 nM D-biotin (Nacalai Tesque, Kyoto, Japan), and 1 μl of the wheat cell-free translational mixture that expressed BirA biotin ligase (~50 ng/μl, BirA: GenBank Accession No. NP_0312927). A 1 × SUB-AMIX solution (125 μl) was placed over the translation mixture. The bilayer was incubated at 26°C for 17 h to allow for protein synthesis. All steps including construction of the DNA templates were performed in the wells of a 96-well plate.

Cleavage assay. The cell-free-synthesized PKs that had luminescent signals > 500 units (in the absence of CASP3) were studied. For each PK, 10 μl of the

CASP3 cleavage buffer (20 mM Tris-HCl, pH 7.5, 0.2 mM DTT, 5 mM MgCl₂, 3 mM ATP, 1 mg/ml BSA, 1 mU CASP3 (Sigma-Aldrich, St. Louis, MO, USA)) was mixed with 1 μl of the translation mixture that contained a Flag-PK-bis ~ biotin construct, and the mixture was incubated at 30°C for 2 h in a well of a 384-well Optiplate (Perkin Elmer, Foster City, CA, USA). Using the reagents of an AlphaScreen IgG (protein A) detection kit (Perkin Elmer) according to the manufacturer's instructions, 15 μl of 20 mM Tris-HCl, pH 7.5, 0.2 mM DTT, 5 mM MgCl₂, 5 μg/ml anti-FLAG M2 antibody (Sigma-Aldrich), 1 mg/ml BSA, 0.1 μl of streptavidin-coated donor beads and 0.1 μl of anti-IgG acceptor beads were added to the well. The solution was incubated at 23°C for 1 h. Luminescence was analyzed using the AlphaScreen detection program (Perkin Elmer). All repetitive mechanical procedures were performed by a Biomek FX robotic workstation (Beckman Coulter, Fullerton, CA, USA). The value of a luminescent signal is reported as the mean of three independent measurements.

TD immunoblotting. A mixture of each Flag-PK-bis ~ biotin construct (3 μl of a translation mixture) and 7 μl of the CASP3 cleavage solution was incubated at 30°C for 1 h in a well of a 384-well Optiplate (Perkin Elmer). Then, the proteins were separated in SDS-PAGE gels and transferred to PVDF membranes (Millipore Bedford, MA, USA). The blotted proteins were prepared for detection using the reagents of an ECL-Plus Western Blotting Detection System kit (GE Healthcare, Piscataway, NJ, USA), anti-Flag M2 antibodies (Sigma-Aldrich) for N-TD, and Alexa488-streptavidin (Invitrogen) for C-TD. The labeled proteins were visualized using a Typhoon Imager (GE Healthcare) with a 532-nm laser and a 526-nm emission filter or an ImageQuant LAS-4000 mini CCD camera system (Fujifilm).

Sequencing and other purification procedures. When possible, long biotinylated C-terminal fragments produced by CASP3 cleavage were recovered attached to streptavidin beads, and then sequenced directly. When a PK construct had a low biotin-labeling efficiency and was cleaved near its C-terminus, a new construct was made by fusing the GST nucleotide sequence encoded in the pEU-E01-Gateway-GST vector to the C-terminal codon of the corresponding PK open-reading frame using the Gateway system and the pEU-E01-Gateway-GST vector. For purification, synthesized PKs (1.2 ml) were purified using Streptavidin Magnesphere Paramagnetic beads (Promega Corp., Madison, WI, USA) for the Flag-PK-bis ~ biotin constructs or glutathione Sepharose 4B (GE Healthcare) for the PK-GST constructs. After washing the beads with PBS, the bound PKs were incubated with CASP3 (15 μl of total volume) as described above. The samples were boiled and the proteins separated by SDS-PAGE. After blotting and visualization (ProBlot, Applied Biosystems, Foster City, CA, USA), the membrane areas that contained the cleaved fragments were cut out and the fragments were sequenced (Applied Biosystems ABI 473A). CSK kinase (Carna Biosciences Inc., Kobe, Japan) and AMPKa1 (Cell Signaling Technology, Beverly, MA, USA) were cleaved with CASP3 (10 μl of total volume), and the cleavage products subjected to MALDI/TOF-MS (Shimazu Techno-Research Inc., Kyoto, Japan) for sequencing. D → A mutagenesis was carried out using the reagents of a PrimeSTAR Mutagenesis Basal kit (TakaraBio, Otsu, Japan) according to the manufacturer's instructions. The mutated genes were sequenced using an ABI PRISM 310 DNA sequencer (Applied Biosystems).

Construction of PK expression plasmids for the cell-based assay. Expression plasmids were produced using the Gateway method. To obtain the attB1-PK-Flag-(stop codon)-attB2 for Gateway BP Clonase II recombination, the open-reading frame products of the 30 newly identified PK substrates of CASP3 that had been produced by PCR using the S1 and T1 primers as described above were PCR amplified using the primers, attB1-S1 (5'-GGGGACAAGTTTGTACAAAAAAGCAGGCTTCACCCACCACCACCA-3') and T1-Flag-stop-attB2 (5'-GGGGACCACCTTTGTACAAGAAAGCTGGGTTTACTTGTGCATCGTCATCCTTG TAGTCGCTTCCAGCACTAGCTCCAGA-3'). These PCR products were each inserted into a pDEST26 vector (Invitrogen) using the Gateway system for construction of the His-PK-Flag nucleotide sequences. All sequences were confirmed by DNA sequencing as described above.

Cell-based assay. HeLa cells were cultured in Dulbecco's modified Eagle's medium, 10% fetal bovine serum, penicillin (100 mg/ml), and streptomycin (50 μg/ml). Transient transfections were carried out using Lipofectamine 2000 Transfection Reagent (Invitrogen) according to the manufacturer's instructions. At 24 h after transfection, cells were harvested after apoptosis was induced. Control cells were treated with DMSO and, for apoptosis induction or inhibition, with 20 ng/ml TNFα

(Calbiochem, La Jolla, CA, USA) and 100 μ M cycloheximide (Chemicon, Temecula, CA, USA) or 125 ng/ml anti-Fas antibody (Medical & Biological Laboratories Co., Ltd., Nagoya, Japan) in the presence (inhibition) or absence (induction) of 100 μ M Z-VAD-FMK (Peptide Institute Inc., Osaka, Japan) for 6 h. Cells were washed with PBS and then lysed directly by adding one volume of 2 \times SDS-PAGE sample buffer (125 mM Tris-HCl, pH 6.8, 20% glycerol, 4% SDS, 10% 2-mercaptoethanol, 0.001% bromophenol blue) before subjecting the cell extracts to SDS-PAGE and immunoblotting, which used anti-His antibodies (GE Healthcare) or anti-Flag M2 antibodies (Sigma-Aldrich). The following antibodies were employed to detect endogenous proteins: anti- α -tubulin (Sigma-Aldrich); anti-AKT2, anti-eEF2K, anti-AMPK α 1, and anti-TGF β R1 (Cell Signaling Technology). Chemiluminescent signals, generated by ECL-Plus reagents (GE Healthcare), or Immobilon Western HRP substrate Luminol Reagent (Millipore), were detected using an LAS-4000 mini biomolecular imager (GE Healthcare).

Conflict of interest

Dr. Endo is a founder of CellFree Sciences Co., Ltd. and a member of its scientific advisory board. Other authors declare no conflict of interest.

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1. Puente XS, Sanchez LM, Overall CM, Lopez-Otin C. Human and mouse proteases: a comparative genomic approach. *Nat Rev Genet* 2003; 4: 544–558.
2. Hunter T. Signaling – 2000 and beyond. *Cell* 2000; 100: 113–127.
3. Manning G, Whyte DB, Martinez R, Hunter T, Sudarsanam S. The protein kinase complement of the human genome. *Science* 2002; 298: 1912–1934.
4. Matrisian LM, Hogan BL. Growth factor-regulated proteases and extracellular matrix remodeling during mammalian development. *Curr Top Dev Biol* 1990; 24: 219–259.
5. Turgeon VL, Houenou LJ. The role of thrombin-like (serine) proteases in the development, plasticity and pathology of the nervous system. *Brain Res Brain Res Rev* 1997; 25: 85–95.
6. van Kempen LC, de Visser KE, Coussens LM. Inflammation, proteases and cancer. *Eur J Cancer* 2006; 42: 728–734.
7. Friedl P, Wolf K. Tube travel: the role of proteases in individual and collective cancer cell invasion. *Cancer Res* 2008; 68: 7247–7249.
8. Abdel-Rahman HM, Kimura T, Hidaka K, Kiso A, Nezami A, Freire E et al. Design of inhibitors against HIV, HTLV-I, and *Plasmodium falciparum* aspartic proteases. *Biol Chem* 2004; 385: 1035–1039.
9. Alnemri ES. Mammalian cell death proteases: a family of highly conserved aspartate specific cysteine proteases. *J Cell Biochem* 1997; 64: 33–42.
10. Hengartner MO. The biochemistry of apoptosis. *Nature* 2000; 407: 770–776.
11. Mahrus S, Trinidad JC, Barkan DT, Sali A, Burlingame AL, Wells JA. Global sequencing of proteolytic cleavage sites in apoptosis by specific labeling of protein N termini. *Cell* 2008; 134: 866–876.
12. Dix MM, Simon GM, Cravatt BF. Global mapping of the topography and magnitude of proteolytic events in apoptosis. *Cell* 2008; 134: 679–691.
13. Righetti PG, Boschetti E. Sherlock Holmes and the proteome – a detective story. *FEBS J* 2007; 274: 897–905.
14. Sawasaki T, Ogasawara T, Morishita R, Endo Y. A cell-free protein synthesis system for high-throughput proteomics. *Proc Natl Acad Sci USA* 2002; 99: 14652–14657.

15. Sawasaki T, Morishita R, Gouda MD, Endo Y. Methods for high-throughput materialization of genetic information based on wheat germ cell-free expression system. *Methods Mol Biol* 2007; 375: 95–106.
16. Takai K, Sawasaki T, Endo Y. Practical cell-free protein synthesis system using purified wheat embryos. *Nat Protoc* 2010; 5: 227–238.
17. Endo Y, Sawasaki T. Cell-free expression systems for eukaryotic protein production. *Curr Opin Biotechnol* 2006; 17: 373–380.
18. Sawasaki T, Hasegawa Y, Morishita R, Seki M, Shinozaki K, Endo Y. Genome-scale, biochemical annotation method based on the wheat germ cell-free protein synthesis system. *Phytochemistry* 2004; 65: 1549–1555.
19. Goshima N, Kawamura Y, Fukumoto A, Miura A, Honma R, Satoh R et al. Human protein factory for converting the transcriptome into an *in vitro*-expressed proteome. *Nat Methods* 2008; 5: 1011–1017.
20. Sawasaki T, Kamura N, Matsunaga S, Saeki M, Tsuchimochi M, Morishita R et al. Arabidopsis HY5 protein functions as a DNA-binding tag for purification and functional immobilization of proteins on agarose/DNA microplate. *FEBS Lett* 2008; 582: 221–228.
21. Takahashi H, Nozawa A, Seki M, Shinozaki K, Endo Y, Sawasaki T. A simple and high-sensitivity method for analysis of ubiquitination and polyubiquitination based on wheat cell-free protein synthesis. *BMC Plant Biol* 2009; 9: 39.
22. Matsuoka K, Komori H, Nose M, Endo Y, Sawasaki T. Simple screening method for autoantigen proteins using the N-terminal biotinylated protein library produced by wheat cell-free synthesis. *J Proteome Res* 2010; 9: 4264–4273.
23. Fischer U, Janicke RU, Schulze-Osthoff K. Many cuts to ruin: a comprehensive update of caspase substrates. *Cell Death Differ* 2003; 10: 76–100.
24. Kurokawa M, Kornbluth S. Caspases and kinases in a death grip. *Cell* 2009; 138: 838–854.
25. Lee KK, Murakawa M, Nishida E, Tsubuki S, Kawashima S, Sakamaki K et al. Proteolytic activation of MST/Krs, STE20-related protein kinase, by caspase during apoptosis. *Oncogene* 1998; 16: 3029–3037.
26. Tomiyoshi G, Horita Y, Nishita M, Ohashi K, Mizuno K. Caspase-mediated cleavage and activation of LIM-kinase 1 and its role in apoptotic membrane blebbing. *Genes Cells* 2004; 9: 591–600.
27. Tatsuta T, Shiraishi A, Mountz JD. The prodomain of caspase-1 enhances Fas-mediated apoptosis through facilitation of caspase-8 activation. *J Biol Chem* 2000; 275: 14248–14254.
28. Packard BZ, Komoriya A, Brotz TM, Henkart PA. Caspase 8 activity in membrane blebs after anti-Fas ligation. *J Immunol* 2001; 167: 5061–5066.
29. Lahti JM, Xiang J, Heath LS, Campana D, Kidd VJ. PITSLRE protein kinase activity is associated with apoptosis. *Mol Cell Biol* 1995; 15: 1–11.
30. Emoto Y, Manome Y, Meinhart G, Kisaki H, Kharbada S, Robertson M et al. Proteolytic activation of protein kinase C delta by an ICE-like protease in apoptotic cells. *EMBO J* 1995; 14: 6148–6156.
31. Casciola-Rosen LA, Anhalt GJ, Rosen A. DNA-dependent protein kinase is one of a subset of autoantigens specifically cleaved early during apoptosis. *J Exp Med* 1995; 182: 1625–1634.
32. Pearce LR, Komander D, Alessi DR. The nuts and bolts of AGC protein kinases. *Nat Rev Mol Cell Biol* 2010; 11: 9–22.
33. Ohoka N, Sakai S, Onozaki K, Nakanishi M, Hayashi H. Anaphase-promoting complex/cyclosome-cdh1 mediates the ubiquitination and degradation of TRB3. *Biochem Biophys Res Commun* 2010; 392: 289–294.
34. Sawasaki T, Hasegawa Y, Tsuchimochi M, Kamura N, Ogasawara T, Kuroita T et al. A bilayer cell-free protein synthesis system for high-throughput screening of gene products. *FEBS Lett* 2002; 514: 102–105.
35. Kamura N, Sawasaki T, Kasahara Y, Takai K, Endo Y. Selection of 5'-untranslated sequences that enhance initiation of translation in a cell-free protein synthesis system from wheat embryos. *Bioorg Med Chem Lett* 2005; 15: 5402–5406.
36. Crooks GE, Hon G, Chandonia JM, Brenner SE. WebLogo: a sequence logo generator. *Genome Res* 2004; 14: 1188–1190.



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Simple Screening Method for Autoantigen Proteins Using the N-Terminal Biotinylated Protein Library Produced by Wheat Cell-Free Synthesis

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Abstract: Autoimmune diseases are a heterogeneous group of diseases characterized by immune reactions against either a major or a limited number of the bodies own autoantigens, causing inflammation and damage to tissues and organs. Thus, identification of autoantigens is an important first step to understanding autoimmune diseases. Here we demonstrate a simple screening method for identification of autoantigens reacting with patient serum antibodies by combination of an N-terminal biotinylated protein library (BPL), produced using a wheat cell-free protein production system, and a commercially available luminescence system. Optimization studies using well-characterized autoantigens showed specific interactions between N-terminal biotinylated proteins and antibody that were sensitively detected under homogeneous reaction conditions. In this optimized assay, 1 μ L of the translation mixture expressing the biotinylated proteins produced significant luminescence signal by addition of diluted serum between 1:500 and 1:10 000 in 25 μ L of reaction volume. For the BPL construction, 214 mouse genes, consisting of 103 well-known autoantigens and 111 genes in the mouse autoimmune susceptibility loci, and the sera of MRL/lpr mouse were used as an autoimmune model. By this screening method, 25 well-known autoantigens and 71 proteins in the loci were identified as autoantigen proteins specifically reacting

with sera antibodies. Cross-referencing with the Gene Ontology Database, 26 and 38 of autoantigen proteins were predicted to have nuclear localization and identified as membrane and/or extracellular proteins. The immune reaction of six randomly selected proteins was confirmed by immunoprecipitation and/or immunoblot analyses. Interestingly, three autoantigen proteins were recognized by immunoprecipitation but not by immunoblot analysis. These results suggest that the BPL-based method could provide a simple system for screening of autoantigen proteins and would help with identification of autoantigen proteins reacting with antibodies that recognize folded proteins, rather than denatured or unfolded forms.

Keywords: autoantigen • autoimmunity • biomarker • cell-free protein production • Gene Ontology • high-throughput screening • MRL/lpr mouse • proteomics

Introduction

Autoimmune diseases are generally characterized by the body's immune responses being directed against its own tissues, causing prolonged inflammation and subsequent tissue destruction.¹ A hallmark of autoimmune diseases is the production of autoantibodies such as antinuclear, anti-Sm and anti-dsDNA in systemic lupus erythematosus (SLE),² and the presence of RF, hnRNP A2 and calpastatin in rheumatoid arthritis (RA).³ However, there are still a lot of autoimmune diseases for which antibodies have not been identified.² To understand the molecular mechanisms in autoimmune diseases, it is important to identify the relevant autoantigens, and moreover, they could be pathogenic in these diseases. It is widely hypothesized that proteins are the major antigenic targets associated with autoimmune diseases.² Therefore, development of methods that allow large-scale screening of autoantigen proteins is indispensable for elucidation and diagnosis of the autoimmune diseases.

To date, autoantigen proteins have been detected as antigenic molecules that are recognized by humoral antibodies, including those in serum.² The large-scale screening of au-

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toantigen proteins reacting with patient serum antibodies has been carried out by mainly three technologies: serological proteome analysis (SERPA), serological expression cloning (SEREX) and protein microarray.⁴ The utility of SERPA and SEREX for this screening is limited because particular cells and tissues are generally used as antigen resources in these systems and they are dependent on artificial membranes for immunoblotting which do not maintain native protein structure.⁵ Recent advances in protein microarray technology have allowed large-scale screening of autoantigens reacting with the sera of patients suffering from autoimmune disorders and cancer.^{5–7} However, protein microarray is not yet a commonly used biochemical tool for screening.⁸ One issue with protein microarrays is that purified recombinant proteins are required, which demonstrate batch-to-batch variation and limited stability and shelf life.⁵ Additionally, it is difficult to maintain the functional form of a protein after their immobilization on a microplate. Many proteins needed to be appropriately oriented for proper functioning.⁹ In fact, a number of spotted autoantigens were not always detectable with planar arrays, presumably due to loss of three-dimensional structures, steric interference or electrostatic repulsion.⁶

In this work, we developed a novel autoantigen protein screening method that overcame the following issues highlighted above: (1) utilization of a high-throughput and genome-wide protein expression system, (2) specific protein labeling for assay using unpurified protein samples and (3) high-throughput detection system of properly folded antigen. Toward the first, we recently developed an automated protein production robot utilizing a high-throughput wheat embryo derived cell-free protein production system.^{10,11} The combination of an automatic cell-free protein production system and the full-length cDNA allowed for facile construction of a robust protein library.¹² To enhance the utility of the library, per the second issue above, specific labeling of each protein is required for efficient detection. We selected biotin as the labeling compound because it is readily available and demonstrates high specificity for streptavidin binding. The biotinylated protein library (BPL) was constructed using target proteins fused to a biotin ligation site (bls), and expression was performed in the presence of biotin and biotin ligase (BirA).¹³ BirA from *Escherichia coli* specifically conjugates a single biotin on the bls. This method was compatible with our high-throughput automated platform. To address the third issue, we selected the luminescent high-throughput protein–protein interaction detection system AlphaScreen.^{14,15} This method can directly recognize biotinylated protein in the translation mixture without purification and the use of any potential protein denaturants allowing for antibody detection of natively folded antigens.¹⁵ In this work, we demonstrate a simple BPL-based method for screening of autoantigen proteins reacting with the sera of an autoimmune disease model mouse, MRL/Mp-*lpr/lpr* (MRL/*lpr*), and the detection of the autoantigen proteins by immunoprecipitation, rather than immunoblotting methods often accompanied by protein denaturation.

Materials and Methods

General. The following procedures have been either described in detail or cited previously:^{10,16} generation of DNA template by polymerase chain reaction (PCR) using “split-primer”; synthesis of mRNA and protein in parallel; estimation of the amounts of synthesized proteins by densitometric scanning of the Coomassie brilliant blue (CBB)-stained bands

or by autoradiography. The wheat germ extract was purchased from Cell-Free Science Co. (Yokohama, Japan). Anti-p53 monoclonal antibody (D01) was purchased from Santa Cruz Biotechnology (Santa Cruz, CA). Mouse serum for mouse immunoglobulin in Figure 1 was purchased from Calbiochem (Darmstadt, Germany). Other reagents used in this study were described previously.^{10,15}

Serum Samples. MRL/*lpr* mice were originally purchased from the Jackson Laboratory (Bar Harbor, ME). All of the mice used in this paper were maintained in clean rooms at the Animal Research Institute, School of Medicine, Ehime University. Sera of female mice were collected from 15 mice and pooled and stocked in -20°C until use. All experiments were done according to the Guidelines for the Care and Use of Laboratory Animals at Ehime University.

DNA Template Construction for the BPL. Functional annotation of mouse (FANTOM) as a mouse full-length cDNA resource is purchased from a company (Danaform, Tokyo, Japan). The DNA templates for transcription were constructed by “split-primer” PCR technique described previous reports.^{10,17} The first PCR was amplified with 10 nM of each of the following primers: a gene specific primer, 5'-CCACCCACCACCACCAAT-Gnnnnnnnnnnnnnnnnnnnnnn (n denotes the coding region of the target gene), and AODA2303 (5'-GTCAGACCCCGTAGAAAAGA) or AODS (5'-TTTCTACGGGGTCTGACGCT). The second PCR products for protein synthesis were constructed with 100 nM SPu 5'-GCGTAGCATTTAGGTGACACT, 1 nM deSP6E02bls-S1 (5'-GGTGACACTATAGAAGTACCTATCTCTCTACACAAAACA-TTCCCTACATACAACCTTTCAACTTCTATTATGGGCCTGAAC-GACATCTTCGAGGCCAGAGATCGAGTGGCAGCAACTCCACCCACCACCACCAATG) and 100 nM AODA2303 or AODS. By this “split-primer” PCR, the bls was fused onto the N-terminals of all the genes for protein biotinylation.¹³

Construction of the BPL by the Cell-Free Protein Synthesis System. Cell-free construction of the BPL is based on the previously described bilayer diffusion system in which 1 μL (50 ng) crude cell-free expressed BirA was added to the translation layer and 500 nM D-biotin (Nacalai Tesque, Kyoto, Japan) was added to both the translation and substrate layers.^{13,18} *In vitro* transcription and cell-free protein synthesis for the BPL were carried out using the GenDecoder1000 robotic synthesizer (CellFree Sciences Co.) as previously described.^{17,19}

Detection of Biotinylated Protein–Antibody Reaction by Luminescence Method. The AlphaScreen assay was performed according to the manufacturer's protocol (PerkinElmer Life and Analytical Sciences, Boston, MA). Reactions were carried out in 25 μL of reaction volume in 384-well Optiwell microtiter plates (PerkinElmer Life and Analytical Sciences). For the antigen–autoantibody reaction, the translation mixture expressing the biotinylated protein was mixed with MRL/*lpr* mouse serum diluted 1:600 in 15 μL of reaction buffer [100 mM Tris-HCl (pH 8.0), 0.01% (v/v) Tween-20 and 0.1% (w/v) bovine serum albumin] and incubated at 26 $^{\circ}\text{C}$ for 30 min. Subsequently, 10 μL of streptavidin-coated donor beads and protein A-conjugated acceptor beads (PerkinElmer Life and Analytical Sciences) were added to a final concentration of 20 $\mu\text{g}/\text{mL}$ per well and incubated at 26 $^{\circ}\text{C}$ for 1 h in a dark box. Fluorescence emission was measured with the EnVision plate reader (PerkinElmer Life and Analytical Sciences), and the resultant data were analyzed using the AlphaScreen detection program. All repetitive mechanical procedures were performed by a Biomek FX robotic workstation (Beckman Coulter, Fullerton, CA).

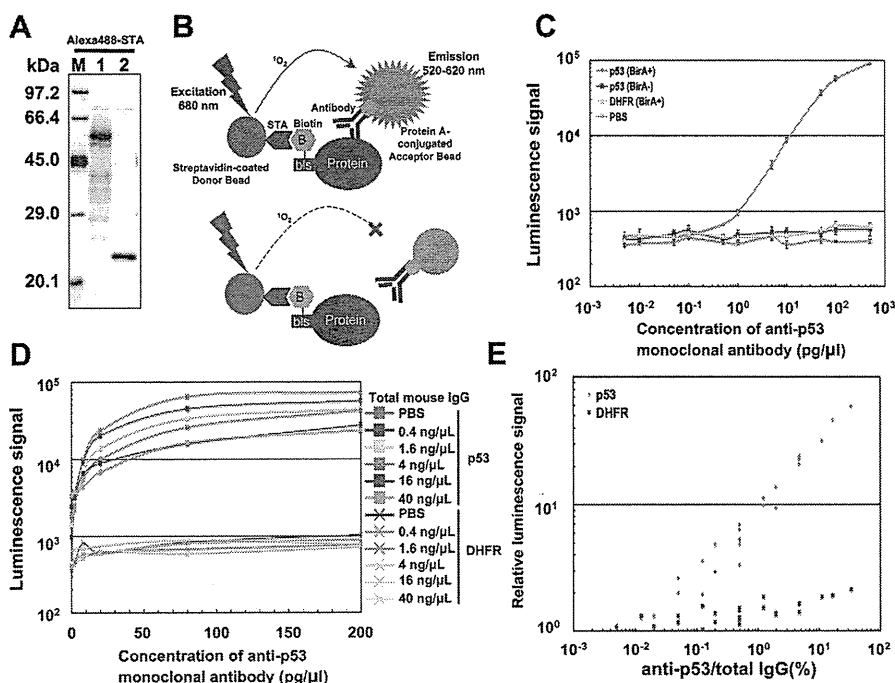


Figure 1. Sensitivity and specificity for detection of biotinylated p53 protein–antibody complex. (A) Biotinylated p53 (lane 1) and dihydrofolate reductase (DHFR) (lane 2) was detected by immunoblotting analysis using Alexa488-STA. M indicates protein molecular weight marker. (B) Schematic diagram of detection of biotinylated protein–antibody interaction by luminescence analysis. When a biotinylated protein and antibody interact (upper panel), Protein A-conjugated acceptor beads bound to antibody and streptavidin (STA)-coated donor beads bound to biotinylated protein are in close proximity. Upon excitation at 680 nm, a singlet oxygen is generated by the donor beads, transferred to the acceptor beads within 200 nm, and the resultant reaction emits light at 520–620 nm. This emission is measured using an EnVision. (C) Detection sensitivity of the antibody concentration measured by luminescence analysis. Translation mixture (1 μ L) expressing biotinylated or nonbiotinylated p53 protein and biotinylated DHFR were incubated with various concentration of monoclonal antibody from 5×10^{-3} to 5×10^2 pg/ μ L. (D) Biotinylated protein–antibody complex by interaction between biotinylated p53 protein and the monoclonal antibody in the presence of mouse serum was detected by luminescence analysis. (E) Minimum IgG amount in the presence of mouse serum to detect biotinylated p53 proteins. The relative luminescence signals between the specific luminescence and background signals indicated in the y-axis.

Immunoblotting. Biotinylated proteins were partially purified using streptavidin-coated beads (Streptavidin Sepharose High Performance, GE Healthcare, Buckinghamshire, U.K.). Translation mixtures (150 μ L) including biotinylated proteins were mixed with 10 μ L of streptavidin-coated beads for 30 min. The resin was washed three times with PBS buffer and then boiled in 15 μ L of SDS sample buffer (50 mM Tris-HCl pH 6.8, 2% SDS, 10% glycerol and 0.2% bromophenol blue). After separation by 12.5% SDS-PAGE, the proteins were transferred to PVDF membrane (Millipore, Bedford, MA) by semidry blotting. The membrane was soaked in PBS containing 5% (w/v) skim milk for 1 h and then incubated with serum diluted 1:200 in PBS containing 0.1% (v/v) Tween 20 (PBST) for 1 h. After washing three times in PBST, it was incubated in PBS including goat-antimouse IgG-HRP antibody (GE Healthcare) diluted 1:10 000 for 30 min. After washing three times in PBST, the blots were detected by the ECL plus detection system (GE Healthcare) by using Typhoon 9400 imaging system (GE Healthcare) according to the manufacturer's protocol.

Immunoprecipitation. Fifty microliters of translation mixture expressing biotinylated proteins were incubated in 50 μ L of IP buffer [PBS containing 0.1% (w/v) BSA, 0.15% (v/v) Tween 20] with 1 μ L of undiluted serum overnight at 4 $^{\circ}$ C. Immobilized Protein A sepharose (20 μ L of 50% slurry, Protein A Sepharose 4 Fast Flow, GE Healthcare) in IP buffer was added to each sample and incubated for 60 min at 4 $^{\circ}$ C. After centrifugation for 1 min at 900 \times g, samples were washed three times with IP

buffer and then boiled for 5 min in SDS sample buffer. After separation by 12.5% SDS-PAGE, the samples were transferred to a Hybond-LFP PVDF membrane (GE Healthcare). After blocking with 5% (w/v) skim milk in PBS overnight at 4 $^{\circ}$ C, the membranes were soaked in PBS buffer containing 10 μ g/mL streptavidin Alexa Fluor 488 conjugate (Alexa488-STA) (Invitrogen, Carlsbad, CA) and were washed three times with PBST. The biotinylated proteins on membrane were detected by Typhoon 9400 imaging system (GE Healthcare) according to the manufacturer's protocol.

Results

Sensitivity and Specificity for Detection of Antigen–Antibody Interaction Using Biotinylated p53 Protein. We adapted that AlphaScreen technology toward detecting interactions between antigen protein and antibody. To validate this technique, we used p53 protein, a well-characterized antigen protein.²⁰ Biotinylated or nonbiotinylated recombinant p53 and biotinylated recombinant dihydrofolate reductase (DHFR), serving as negative control, were synthesized by the wheat cell-free system (Figure 1A). For the analysis of antigen protein–antibody interaction, the translation mixture was used without any purification. In the AlphaScreen system, interaction of the biotinylated protein and antibody in sera results in a biotinylated protein–antibody complex that is captured simultaneously by the streptavidin-coated donor beads and the protein

A-conjugated acceptor bead. The resultant proximity of the acceptor and donor bead generates the luminescent signal upon irradiation at 680 nm. This is illustrated in Figure 1B.

For biotinylation of the target protein, the N-terminus of the target was fused to the bls, and the cell-free system was supplemented with BirA and biotin.¹³ This biotin ligation method yields a biotin labeling on the bls, indicating a specific recognition of the target protein by AlphaScreen. To investigate the specificity and sensitivity of the antibody detection, translation mixtures expressing biotinylated or nonbiotinylated p53 protein were incubated with various concentrations of monoclonal antibody, ranging from 5×10^{-3} to 5×10^2 pg/ μ L. This luminescence method specifically detected interaction of monoclonal antibody and the biotinylated p53 from the unpurified translation mixture, whereas nonbiotinylated p53 and biotinylated DHFR did not produce a significant luminescent signal (Figure 1C). In this condition, the biotinylated p53 was detected by anti-p53 antibody at concentrations as low as 0.5 pg/ μ L. Next, we investigated whether this luminescence method could detect the biotinylated protein-antibody complex in the presence of mouse serum. Translation mixture expressing biotinylated p53 protein was incubated with various concentrations of monoclonal antibody from 2 to 200 pg/ μ L and mouse immunoglobulin from 0.4 to 40 ng/ μ L. Figure 1D showed that this method could specifically detect the immunocomplex of biotinylated p53 protein and monoclonal antibody in the presence of an excess of mouse immunoglobulin. In addition, comparison between the specific luminescence and background signals indicated that biotinylated p53 could be detected at more than 0.05% of anti-p53 antibody in mouse serum (Figure 1E). These results indicate that this system is a highly specific and sensitive method for detection of interaction between biotinylated recombinant protein and antibody in whole serum.

Detection of Autoantibodies against Hars and LmnB2 Proteins in sera of MRL/lpr Mice. We next tested our protocol for the well-characterized autoantigens histidyl-tRNA synthetase (Hars)²¹ and lamin B2 (LmnB2)²² in the autoimmune disease model mouse MRL/lpr.²³ To determine the assay conditions using serum samples, biotinylated recombinant Hars and LmnB2 proteins were used (Figure 2A) to detect autoantibody in the sera of MRL/lpr mice. Cell-free synthesis of biotinylated Hars and LmnB2 demonstrated yields of 820 and 600 nM, and 43.0 and 56.4% of biotinylation, indicating biotinylated Hars and LmnB2 proteins were 354.4 and 338.5 nM, respectively. Various volumes (0.003 to 4 μ L) of translation mixture expressing biotinylated Hars or LmnB2 protein were incubated with the serum of MRL/lpr mouse (final 1:1000 dilution) in 25 μ L of reaction volume (Figure 2B). Significant luminescent signals were observed at additions of biotinylated Hars or LmnB2 proteins between 0.01 and 1 μ L, which corresponds to biotinylated protein concentrations between 0.14 and 14 nM or 0.13 and 13 nM, respectively. Also serum dilutions between 1:500 and 1:10 000 produced high luminescence signal in 25 μ L of reaction volume using 1 μ L of the translation mixtures (Figure 2C). These results mean that five micro litter of serum and 200 μ L of cell-free translation mixture expressing biotinylated proteins would be sufficient for 200 assays. Taken together, these results suggest that the luminescence method using cell-free expressed biotinylated proteins

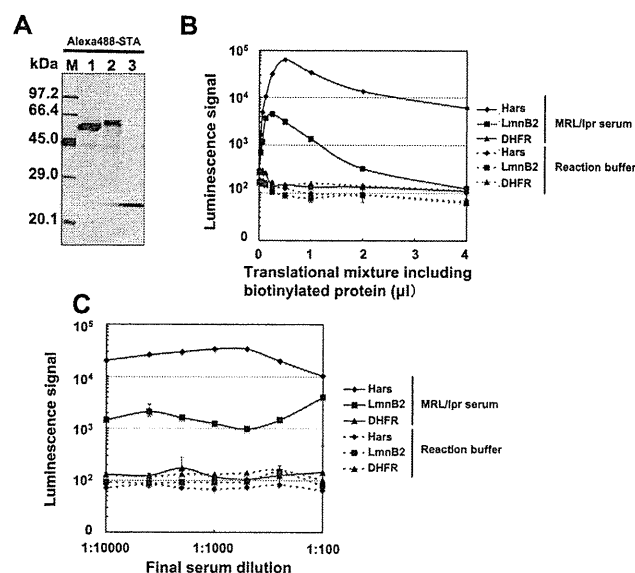


Figure 2. Detection of autoantibodies against Hars and LmnB2 proteins. (A) Biotinylated histidyl-tRNA synthetase (Hars) (lane 1), and lamin B2 (LmnB2) (lane 2) and DHFR (lane 3) proteins were detected by immunoblotting analysis using Alexa488-STA. M indicates protein molecular weight marker. (B) Various volumes (0.003 to 4 μ L; representing 2.5–3280 nM Hars, 1.8–2400 nM LmnB2 and 6.4–8520 nM DHFR) of translation mixture expressing biotinylated Hars, LmnB2 or DHFR proteins were incubated with serum of MRL/lpr mouse sera (final 1:1000 dilution) in 25 μ L of reaction volume. (C) Serum dilution between 1:100 and 1:10 000 was incubated with 1 μ L of the translation mixtures in 25 μ L of reaction volume.

would be useful for screening the reaction of autoantigen proteins with autoantibodies in serum.

Construction of the BPL by the Wheat Cell-Free Protein Production System. It has long been thought that comprehensive screening using a protein library is a strong tool for identification of antigen proteins.^{12,24,25} The scheme for the BPL-based screening is shown in Figure 3A. To construct the N-terminal BPL, we selected 226 genes (Supplementary Table 1, Supporting Information) that included well-known autoantigen proteins and proteins coded by genes in the mouse autoimmune susceptibility loci²⁶ from the mouse full-length cDNA resource (FANTOM).^{27,28} For biotinylation, a bls was fused onto 5' site of a target gene by "split-primer" PCR.¹³ Using the PCR, 222 (98.2%) out of 226 genes were successfully amplified and of those, 217 (96%) were transcribed. Synthesis of biotinylated proteins was performed on the GenDecoder1000,¹⁹ and expression confirmed by SDS-PAGE combined with immunoblot analysis using Alexa488-STA (Figure 3B). Finally 214 clones (94.6%) were produced as biotinylated proteins (Supplementary Table 1, Supporting Information) at maximum and minimum concentrations of 500 and 10 nM respectively (data not shown). From our results in Figure 2B, the immunoresponse of biotinylated proteins could be detected below 0.2 nM by the luminescence method, indicating that all 214 proteins are at concentrations viable for screening. Therefore, we used these proteins as the BPL for screening of autoantigen proteins.

BPL-Based Screening of Autoantigen Proteins Using the MRL/lpr Mouse Sera. To identify autoantigen proteins reacting with antibodies in serum of autoimmune disease mice, the BPL and sera from pools of MRL/lpr or normal mouse sera (NMS)

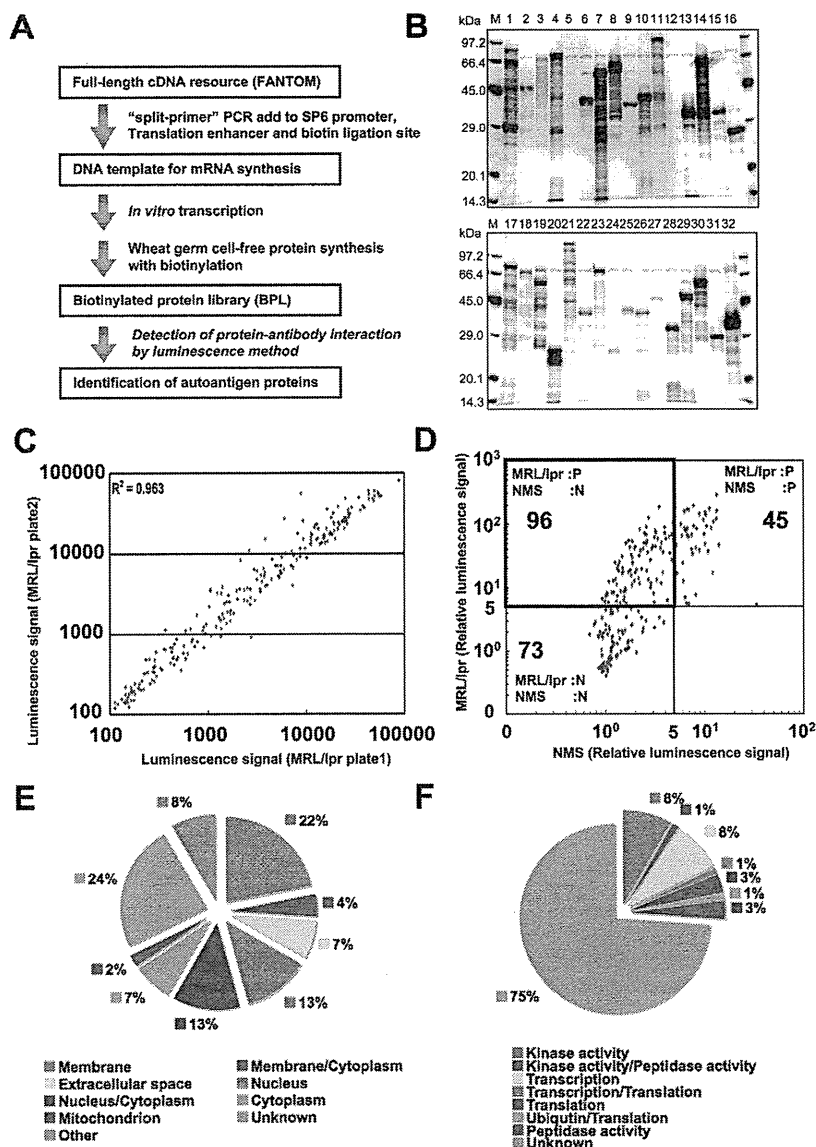


Figure 3. BPL-based screening of autoantigen proteins using the MRL/lpr mouse sera. (A) Schematic of the BPL-based screening method. (B) Thirty-two randomly selected biotinylated proteins of the BPL were detected by immunoblotting analysis using Alexa488-STA. (C) Scatter plot showing the luminescent signals in each well of two independent screening data sets using MRL/lpr mouse sera. The x-axis indicates luminescence signals in MRL/lpr plate 1 whereas the y-axis represents those in MRL/lpr plate 2. (D) Each data point represents luminescence signals using MRL/lpr mouse sera or normal mouse sera (NMS). The x-axis indicates luminescence signals in NMS whereas the y-axis represents those in MRL/lpr mice. (E, F) Ninety-six proteins identified as autoantigen proteins were grouped by protein localization in cells (E) Membrane (GO:0016020), Nucleus (GO:0005634), Cytoplasm (GO:0005737), Extracellular region (GO:0005576) and Mitochondrion (GO:0005739) and biological function/process (F) Kinase activity (GO:0016301), Peptidase activity (GO:0008233), Ubiquitin (GO:0005551), Translation (GO:0006412) and Transcription (GO:0006350) according to Gene Ontology Database. Minor groups less than 3 proteins were belonged to "Other" group. More detailed information on individual proteins was indicated in the Table 1.

were used. In each well of a 384-well plate, a translation mixture expressing biotinylated protein was incubated with either serum for 30 min, and subsequently a mixture of donor and acceptor beads was added to each well (see Figure 1B). After incubation, antigen-antibody reaction of the BPL was detected by the luminescence assay as described above. As shown in the scatter plot (Figure 3C), the intensity of paired luminescent signals in each well of two independent screening data sets (plate 1 and 2) using MRL/lpr mouse sera showed a linear distribution with a R^2 of 0.963, indicating reproducibly sufficient score for screening. We then compared the luminescent signals of the BPL reacted with MRL/lpr sera and NMS (Figure 3D). In

this assay, only 141 of the 214 proteins in the BPL were identified as positive clones, which was indicated by a luminescence signal 5-fold higher than the average background signal. Only 96 proteins in the 141 positive clones reacted with MRL/lpr sera, whereas the remaining 45 proteins interacted with both sera. From these results, 96 proteins were identified as autoantigen proteins in MRL/lpr mice (Table 1, upper left panel in Figure 3D). In these MRL/lpr autoantigen proteins, 25 well-known autoantigens were included, and 71 out of 96 clones were coded in the genetic loci on chromosome 10; 40 cM, chromosome 15; 18 cM and chromosome 19; 49 cM,²⁶ indicating that this screening identified new MRL/lpr sera