# Genome-wide analysis to predict protein sequence variations that change phosphorylation sites or their corresponding kinases

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#### **ABSTRACT**

We define phosphovariants as genetic variations that change phosphorylation sites or their interacting kinases. Considering the essential role of phosphorylation in protein functions, it is highly likely that phosphovariants change protein functions. Therefore, a comparison of phosphovariants between individuals or between species can give clues about phenotypic differences. We categorized phosphovariants into three subtypes and developed a system that predicts them. Our method can be used to screen important polymorphisms and help to identify the mechanisms of genetic diseases.

#### INTRODUCTION

Protein phosphorylation is involved in various important processes: development and learning at the organism level, and the cell cycle, differentiation and apoptosis at the cellular level (1,2). Phosphorylation can change the subcellular localization of a protein, its life span and its affinity for other proteins or DNA (3). Therefore, the addition or deletion of phosphorylation sites through phosphovariants can lead to functional variations in proteins that can result in phenotypic variations or genetic diseases. By our definition, phosphovariants are variations that change phosphorylation sites or their interacting kinases. We propose three subtypes of phosphovariants. First, some variations occur directly at phosphorylation sites, and these sites will be removed if the phosphoreceptors are replaced with amino acids other than serine, threonine or tyrosine. Conversely, replacement of another amino acid with a serine, threonine or tyrosine may add a new

phosphorylation site. Second, variations adjacent to phosphorylation sites can result in the removal or addition of phosphorylation sites. Third, variations may change the kinases that recognize phosphorylation sites, without changing the phosphorylation site itself. We divided phosphovariants into type I, II and III, respectively, according to the above descriptions (Figure 1).

We developed PredPhospho (version 2), a web-based computer program that predicts phosphorylation sites, and PhosphoVariant, a database for human phosphovariants. Even the advanced laboratory techniques used to analyze phosphorylation sites, such as mass spectrometry (MS), cannot analyze all types of proteins (4,5). For example, peptides that are either too small or too large in mass can be easily missed. Moreover, membrane proteins cannot be obtained in sufficient quantities for analysis (5). Even when proteins can be analyzed with MS, it is very time consuming and expensive to make thousands of variant proteins and select the phosphovariants. PredPhospho can predict the phosphorylation sites in kinase-specific ways, using the support vector machines (SVMs) derived from statistical learning theory proposed by Vapnik and Chervonenkis in 1995 (6). In our study, we searched for known phosphovariants and tried to predict other possible phosphovariants among human variations.

# **METHODS**

## **PredPhospho**

We created classifiers of various kinases by training SVMs with phosphorylation site sequences and nonphosphorylated site sequences. In other words, our classifiers determine whether serine, threonine or tyrosine residues within a sequence can be phosphorylated or not. 'Phosphorylated site sequences' refers to peptide sequences with a serine,

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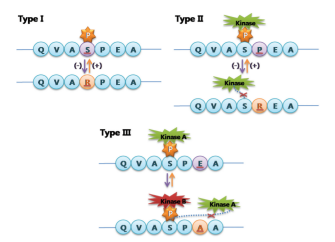


Figure 1. Schematic illustration of phosphovariants according to their

threonine or tyrosine residue located at the center, and which are phosphorylated. Conversely, 'nonphosphorylated site sequences' are sequences with a serine, threonine or tyrosine residue located at the center, which have not been found, yet, to be phosphorylated. We obtained phosphorylated site sequences from public databases: the Swiss-Prot (release 54.8) and the Human Protein Resource Database (HPRD, release 7). Nonphosphorylated site sequences were taken from laboratory data confirmed by MS (see Supplementary Data).

Manning et al. (7) found 518 human protein kinase genes in the human genome sequence, using the hidden Markov model (HMM) profile, and confirmed the identities of more than 90% of the identified kinase genes using cDNA cloning. They also classified the protein kinase superfamily into nine broad groups, and subdivided the groups into 134 families and 204 subfamilies, using sequence comparisons of the kinase catalytic domains. We classified the phosphorylated site sequences according to their kinases and created the classifiers in a kinasespecific manner. Because of the limitations of the phosphorylated sequence data presently available in public databases, we can make classifiers for only six kinase groups: AGC, CAMK, CK1, CMGC, STE and TK; and 18 kinase families: AKT, CAMK2, CAMKL, CDK, CK1, CK2, GSK, IKK, JakA, MAPK, PDGFR, PIKK, PKA, PKC, RSK, Src, STE20 and Syk (all abbreviations are shown in the footnote to Supplementary Table S1). The detailed algorithms and methods were described in the Supplementary Data.

#### **Evaluation of the system**

The performance status of the prediction for each kinase group and family is shown in Supplementary Table S2. The performance of the prediction with combinations of all the kinase group models or all the family group models is not the numerical multiplication of the performance of each model. Therefore, to evaluate the performances of the predictions at the kinase group level or at the family level, we tested two proven real data sets, which were

compiled with MS experiments. Data set I was created by Olsen et al. (5), who identified phosphorylation sites in proteins from HeLa cells and classified the phosphorylation sites according to their definition. Four classes are based on their localization probabilities: <0.25, 0.25 < < 0.75 without kinase motifs, 0.25 < < 0.75 with kinase motifs and  $\geq 0.75$ . We used only monophosphopeptides that had localization probabilities of at least 0.75. Data set II was derived from the paper of Beausoleil et al. (4) and we used those of their phosphopeptides with a localization certainty of >99.4% (see Supplementary Figure S2). We selected phosphorylation sites and nonphosphorylated sites from these two data sets. To avoid overestimating the performance of PredPhospho, we discarded sequences that were more than 70% identical to sequences used for training PredPhospho to sequences used for training PredPhospho (Table 4). We also avoided using false nonphosphorylated sites by omitting those sites that are listed as phosphorylation sites in Swiss-Prot or HPRD. We tested the two kinds of data sets not only with our PredPhospho, but also with Scansite.

#### **Prediction of phosphovariants**

We extracted information about human genetic variations from SwissVariant of the Swiss-Prot database. SwissVariant includes single amino acid polymorphisms and missense mutations (8). The number of variations listed in SwissVariant was 33651. We consulted the Swiss-Prot database and HPRD about their effects, and the references to these variations and phosphorylation sites. With PredPhospho and Scansite, we predicted the phosphorylation sites and related kinases for the original sequences and the variant sequences. The phosphovariants could be identified when the phosphorylation sites or interacting kinases were altered between the original sequence and the variant sequence. If the phosphorylation site is in the same location as the variation, it is type I. In type II phosphovariants, the variation is not in the same location as the phosphorylation site. We added the symbol (+) to types I and II when the phosphovariants added new phosphorylation sites, and (-) when the phosphovariants removed phosphorylation sites [e.g. type I (+) or type I (-)]. Type III phosphovariants are caused by changes in the types of kinases involved, rather than in the phosphorylation site itself, regardless of the locations of the variations. One variation can include more than one class of phosphovariant, because one variation can affect two or more phosphorylation sites. We predicted phosphovariants at the kinase group level and the family level. The predictions at the family level are more sensitive, but less specific, than those at the group level. To minimize false negatives, we varied the specificity options (95%, 97%, 98% or 99%) according to the specificity of each model (Supplementary Table S3). The specificity options are described in the Supplementary Data.

#### Sequence logos

We obtained 562 phosphorylation site sequences recognized by the CMGC kinase group from Swiss-Prot and HPRD. We trimmed the sequences as six symmetric residues centered phospohrylation sites. We aligned the sequences and obtained a sequence logo using the web program (http://weblogo.berkeley.edu/logo.cgi).

#### WWW programs

The PredPhospho (version 2) and PhosphoVariant were implemented using the PERL (version 5.8.8) programming language and MySQL (version 5.0.18). The PhosphoVariant is a database for the definite and possible human variants changing phosphorylation sites and their interacting kinases. They are available at: http://phospho variant.ngri.go.kr/seq input predphospho2.htm http://phosphovariant.ngri.go.kr, respectively.

Bug fixing and minor changes of database will be done, whenever needed. The new version of Swiss-Prot and HPRD will be incorporated into our database, annually at least.

#### **RESULTS**

# Type I phosphovariants

The substitution of phosphoreceptor amino acids with amino acids other than serine, threonine or tyrosine causes the elimination of phosphorylation sites and can be classified as type I (-) phosphovariants according to our classification. We found 50 type I phosphovariants by matching the locations of the variations and those of phosphorylation sites registered in the Swiss-Prot database and the Human Protein Resource Database (HPRB, Table 1) (9,10). Of these phosphovariants, 19 are known to cause Mendelian-inherited diseases and 18 are associated with cancers. Another 13 phosphovariants are polymorphisms.

Conversely, new phosphorylation sites can be created by variations and we defined these as type I (+) phosphovariants. We found an example in the study of Nousiainen et al. (11) The cell line that they used had a Gly766Ser mutation in the probable ATP-dependent RNA helicase DDX27 (Swiss-Prot ID, O96GO7), which was identified as phosphorylated. Similarly, the polymorphisms in Tables 1–3 are good examples of the addition of phosphorylation sites by sequence variations. For example, if we postulate that the isoleucine at amino acid 571 of CTP synthase 1 (Swiss-Prot ID, P17812) is changed to serine, then it also represents a type I (+) phosphovariant, rather than a type I (–) phosphovariant (Table 1).

# Type II phosphovariants

It is more difficult to find examples of type II phosphovariants than to find type I phosphovariants, because we cannot definitely say that a phosphorylation site is changed by a substitution near a phosphorylation site. However, when some kinases (although not all kinases) recognize phosphorylation sites, the specific amino acids near the phosphoreceptor are important. In Figure 2, we present sequence logos of phosphorylation site sequences for the CMGC kinase group. The proline residues at position +1 relative to the phosphorylation sites are important in the phosphorylation site sequences of the CMGC kinase group, especially the CDK kinase family and the MAPK kinase family. Most (84%) of the phosphorylation sites of the CMGC group registered in the Swiss-Prot database and the Human Protein Resource Database (91% of the CDK and 87% of the MAPK kinase families) have +1 proline residues. If the proline is substituted with another amino acid, it is highly probable that the adjacent phosphorylation site will be abolished. The phosphorylation site at Ser112 of peroxisome proliferatoractivated receptor gamma protein (Swiss-Prot ID, P37231) is eliminated by the Pro113Gln substitution (12). We found three other polymorphisms that abolish phosphorylation sites of the CMGC kinase group, but the removal of these phosphorylation sites has not yet been confirmed (Table 2). The presence of specific amino acids does not directly affect phosphorylation by kinases other than those of the CMGC kinase group, but sequences near the phosphorylation site must be considered. Kinases recognize the residues surrounding the target phosphorylation site, and the amino acids bordering phosphorylation sites are, in turn, affected by other nearby residues (13). Hence, when the relevant kinases are not members of the CMGC kinase group, it is difficult to predict type II phosphovariants simply by database matching, without specific programs that predict phosphorylation sites.

# Type III phosphovariants

Type III phosphovariants are those variations that change only the type of kinase involved, without affecting the phosphorylation site itself. For example, Ser386 of the tyrosine protein phosphatase, nonreceptor type 1 (PTPN1, Swiss-Prot ID, P18031) is phosphorylated by cell division cycle 2 (CDC2) kinase, a member of the CMGC kinase group, and by casein kinase 2 (CK2), a member of the 'Other' kinase group (14,15). The Pro387Leu substitution reduces 75% of the phosphorylation by CDC2 in vitro (16). However, it has not been confirmed that Pro387Leu inhibits the recognition of Ser386 by CK2. Only about 5% of the sites phosphorylated by CK2 that are registered in the Swiss-Prot database or HPRD have a proline residue at position +1 relative to the phosphorylation site. There is also no known consensus sequence for CK2 that contains proline at that location (17). Therefore, we infer that the proline residue is not essential for phosphorylation by CK2 and that Pro387Leu will have little effect on phosphorylation by CK2. Therefore, we consider Pro387Leu of PTPN1 a type III phosphovariant because it inhibits the recognition of Ser386 by CDC2 kinase but has little effect on its phosphorylation by CK2 (16,18).

Kinases that recognize serine and threonine differ from the kinases that recognize tyrosine. The substitution of phosphorylated serine or threonine for tyrosine, or vice versa, can remove a phosphorylation site or change the type of kinase that recognizes it. Changes between serine and threonine can also cause changes in the phosphorylation site and the responsive kinase. Therefore, the phosphovariants in Table 3 (Panel b) are either type I or type III phosphovariants.

Table 1. Examples of type I phosphovariants

Gene name (Swiss-Prot ID)	Variation site <sup>a</sup> (Swiss-Prot variant ID)	Phosphory- lation site	Local peptide sequence <sup>b</sup>	Effect <sup>c</sup>	Reference(s) for variation <sup>d</sup>	Reference(s) for phosphor- ylation site <sup>e</sup>
Panel a: Type I(-) p	hosphovariants					
Phosphovariants causi	ing Mendelian inherited di	sease				
EDNRB (P24530) FANCA (O15360)	S305N (VAR_003472) S858R (VAR_017498)	S305 S858	CEMLRKK S GMQIALN QSRDTLC S CLSPGLI	Hirschsprung disease type 2 Fanconi anemia	8852659 10094191 11091222	14636059 17924679
KCNJ1 (P48048) L1CAM (P32004)	S219R (VAR_019726) S1194L (VAR_003947)	S219 S1194	RVANLRK S LLIGSHI AFGSSQP S LNGDIKP	Bartter syndrome type 2 Hydrocephalus due to stenosis of the aqueduct of Sylvius mental retardation, aphasia, shuffling gait and adducted	8841184 8556302 7881431	8621594 17081983
MAPT (P10636)	S622N (VAR_010350)	S622	KHVPGGG S VQIVYKP	thumbs syndrome Frontotemporal dementia and parkinsonism linked	10208578	7706316
MAPT (P10636)	S637F (VAR_019665)	S637	VDLSKVT S KCGSLGN	chromosome 17 Pick disease	11891833	11104762 9199504
MAPT (P10636)	S669L (VAR_019667)	S669	DFKDRVQ S KIGSLDN	Fatal respiratory	14595660	11104762
MITF (O75030)	S405P (VAR_010302)	S405	QARAHGL S LIPSTGL	hypoventilation Waardenburg syndrome type IIa	8589691	10587587
NFKBIA (P25963)	S32I (VAR_034871)	S32	LLDDRHD S GLDSMKD		14523047	10882136 9721103 8601309 16319058 10723127
PER2 (O15055)	S662G (VAR_029080)	S662	ALPGKAE S VASLTSQ	Familial advanced sleep-phase syndrome	11232563	9214631 11232563
PTPN11 (Q06124)	Y62D (VAR_015605)	Y62	KIQNTGD Y YDLYGGE	Patients with Noonan syndrome 1 manifesting juvenile myelomonocytic leukemia	11992261 12325025 12960218	15951569 15592455
RAF1 (P04049)	S259F (VAR_037809)	S259	SQRQRST S TPNVHMV	Noonan syndrome type 5	12717436 17603483	8349614 11997508 11971957 10576742
RAF1 (P04049) RAF1 (P04049) RPS6KA3 (P51812) STAT3 (P40763)	T491R (VAR_037819) T491I (VAR_037818) S227A (VAR_006195) Y657C (VAR_037381)	T491 T491 S227 Y657	IGDFGLA T VKSRWSG IGDFGLA T VKSRWSG DHEKKAY S FCGTVEY FAEIIMG Y KIMDATN	Noonan syndrome type 5 Noonan syndrome type 5 Coffin–Lowry syndrome Hyperimmunoglobulin E recurrent infection syndrome autosomal dominant	17603483 17603483 8955270 17881745	11447113 11447113 17192257 15037656
TGFBR2 (P37173)	Y336N (VAR_022352)	Y336	AKGNLQE Y LTRHVIS	Loeys–Dietz aortic aneurysm syndrome	15731757	9169454
TNNI3 (P19429) TSC1 (Q92574)	S166F (VAR_029454) T417I (VAR_009403)	S166 T417	LGARAKE S LDLRAHL SLPQATV T PPRKEER	Hypertrophic cardiomyopathy Tuberous sclerosis complex, could be a polymorphism	12974739 10570911 10607950	11121119 14551205
Phosphovariants found	d in cancer					
CDH1 (P12830) CTNNB1 (P35222)	S838G (VAR_001322) S23R (VAR_017612)	S838 S23	LVFDYEG S GSEAASL PDRKAAV S HWQQQSY	Ovarian cancer Hepatocellular carcinoma,	8075649 10435629	10671552 12027456
CTNNB1 (P35222)	S33F (VAR_017617)	S33	QQQSYLD S GIHSGAT	no effect Pilomatrixoma, medulloblastoma and	12027456 10666372 10435629	12000790 12114015
CTNNB1 (P35222)	S33L (VAR_017618)	S33	QQQSYLD S GIHSGAT	hepatocellular carcinoma Hepatocellular carcinoma	10192393 10435629	11818547 12000790 12114015
CTNNB1 (P35222)	S37A (VAR_017624)	S37	YLDSGIH S GATTTAP	Medulloblastoma, hepatocellular carcinoma	12027456 10435629 10666372	11818547 12000790 12114015 11818547
CTNNB1 (P35222)	S37C (VAR_017625)	S37	YLDSGIH S GATTTAP	Pilomatrixoma, hepatoblastoma	9927029 10192393	12000790 12114015
CTNNB1 (P35222)	S37F (VAR_017626)	S37	YLDSGIH S GATTTAP	Pilomatrixoma	10192393	11818547 12000790 12114015 11818547

(continued)

Table 1. Continued

Gene name (Swiss-Prot ID)	Variation site <sup>a</sup> (Swiss-Prot variant ID)	Phosphory- lation site	Local peptide sequence <sup>b</sup>	Effect <sup>c</sup>	Reference(s) for variation <sup>d</sup>	Reference(s) for phosphor- ylation site <sup>e</sup>
CTNNB1 (P35222)	T41A (VAR_017629)	T41	GIHSGAT T TAPSLSG	Hepatoblastoma and hepatocellular carcinoma, also in a desmoid tumor	12051714 10398436 9927029 12027456 10655994 10435629	12051714 12114015 11818547 12000790
CTNNB1 (P35222)	T41I (VAR_017630)	T41	GIHSGAT T TAPSLSG	Pilomatrixoma and hepatocellular carcinoma	10435629 10192393 10435629	12051714 12114015 11818547
CTNNB1 (P35222)	S45F (VAR_017631)	S45	GIHSGAT T TAPSLSG	Hepatocellular carcinoma	10435629	12000790 12051714 12000790 11955436
CTNNB1 (P35222)	S45P (VAR_017632)	S45	GATTTAP S LSGKGNP	Hepatocellular carcinoma	10435629	12051714 12000790 11955436
FAM10A4 (Q8IZP2)	S71L (VAR_023644)	S71	DLKADEP S SEESDLE	B-cell leukemia, multiple myeloma, and prostate cancer	12079276	17081983
MET (P08581)	Y1230C (VAR_006292)	Y1230	FGLARDM Y DKEYYSV		9140397	12475979
MET (P08581)	Y1230H (VAR_006293)	Y1230	FGLARDM Y DKEYYSV		9140397	12475979
NME1 (P15531) RB1 (P06400)	S120G (VAR_004625) S567L (VAR_005579)	S120 S567	GRNIIHG S DSVESAE SLAWLSD S PLFDLIK	Neuroblastoma Retinoblastoma	8047138 10671068 2594029	8810265 10207050
TP53 (P04637)	T155A (VAR_005901)	T155	DSTPPPG T RVRAMAI	Esophageal cancer	1868473	12628923
Phosphovariants relate	ed with polymorphism					
BARD1 (Q99728) C10orf11 (Q9H2I8) CTNND1 (O60716)	S186G (VAR_038371) S153F (VAR_033686) Y217C (VAR_020929)	S186 S153 Y217	SYEFVSP S PPADVSE SSEDVAS S PERHYTP PDGYSRH Y EDGYPGG	Polymorphism (rs16852741) Polymorphism (rs35349706) Polymorphism (rs11570194)		15855157 16964243 15592455 16212419
CTPS (P17812)	S571I (VAR_027055)	S571	RDTYSDR S GSSSPDS	Polymorphism (rs17856308)	15489334	16097034 17081983
HIF1A (Q16665) INSR (P06213) KRT36 (O76013) MYH15 (Q9Y2K3) PDLIM5 (Q96HC4)	T796A (VAR_015854) Y1361C (VAR_015933) T315M (VAR_020306) T1125A (VAR_030238) S136F (VAR_023779)	T796 Y1361 T315 T1125 S136	ESGLPQL T SYDCEVN SYEEHIP Y THMNGGK EIIELRR T VNALEIE KTVKELQ T QIKDLKE PRPFGSV S SPKVTSI	Polymorphism (rs1802821) Polymorphism (rs13306449) Polymorphism (rs2301354) Polymorphism (rs3900940) Polymorphism (rs2452600)	7657032	17382325 11401470 17081983 17081983 17287340
PNN (Q9H307) SUB1 (P53999)	S671G (VAR_023368) S11G (VAR_032870)	S671 S11	HKSSKGG S SRDTKGS SKELVSS S SSGSDSD	Polymorphism (rs13021) Polymorphism (rs17850527)	10095061 15489334	17287340 17081983 16689930
SRRM2 (Q9UQ35) TP53 (P04637)	S883C (VAR_027260) S366A (VAR_022317)	S883 S366	SPDPELK S RTPSRHS PGGSRAH S SHLKSKK	Polymorphism (rs17136053) Polymorphism		17287340 9183006
Panel b: Type I(+) p	hosphovariants					
DDX27 (Q96GQ7)	G766S	S766	ALKQYRA G PSFEERK	Unknown	16565220	16565220

<sup>&</sup>lt;sup>a</sup>Locations and amino acid changes of the variations in the proteins.

<sup>&</sup>lt;sup>b</sup>Peptide sequences with 15-mer amino acids. The amino acids in the eighth position are phosphorylated residues.

The meanings or consequences of the variations. We referred to the feature tables of Swiss-Prot for these effects. If the polymorphisms are enrolled in dbSNP, the IDs of dbSNP are written in the parentheses.

<sup>&</sup>lt;sup>d</sup>Pubmed ID for the references of the variations

ePubmed ID for the references of the phosphorylation sites

Protein names which are abbreviated by their gene names: epithelial cadherin (precursor), CDH1; catenin β-1, CTNNB1; probable ATP-dependent RNA helicase DDX27, DDX27; endothelin B receptor (precursor), EDNRB; protein FAM10A4, FAM10A4; Fanconi anemia group A protein, FANCA; ATPsensitive inward rectifier potassium channel 1, KCNJ1; keratin, type I cuticular Ha6, KRT36; Neural cell adhesion molecule L1, L1CAM; microtubuleassociated protein tau, MAPT; hepatocyte growth factor receptor (precursor), MET; microphthalmia-associated transcription factor, MITF; NF-κ-B inhibitor α, NFKBIA; nucleoside diphosphate kinase A, NME1; period circadian protein homolog 2, PER2; tyrosine-protein phosphatase nonreceptor type 11, PTPN11; RAF proto-oncogene serine/threonine-protein kinase, RAF1; retinoblastoma-associated protein, RB1; ribosomal protein S6 kinase alpha-3, RPS6KA3; signal transducer and activator of transcription 3, STAT3; TGF-beta receptor type-2 (precursor), TGFBR2; cardiac troponin I, TNNI3; cellular tumor antigen p53, TP53; Hamartin, TSC1.

# Performance of PredPhospho and Scansite

We developed prediction models for six kinase groups and 18 kinase families. Their accuracy ranged from 70.80 to 94.67% at the kinase family level and from 71.77 to 91.18% at the kinase group level (Supplementary Table S2). We tested our prediction models using two real laboratory data sets compiled

with MS. For the six kinase group models, the sensitivities were 79.40% with data set I and 75.47% with data set II, but the specificities were as low as 60.62% for data set I and 61.04% for data set II because of the accumulation of false negatives by all six kinase group models. When we modified the specificity to >95% (see Supplementary Data for the modification of the specificity for each model), the specificities increased to 72.09% and

**Table 2.** Examples of type II(-) phosphovariants

Gene name (Swiss-Prot ID)	Variation site (Swiss- Prot variant ID)	Removed phosphorylation site (related kinases)	Local peptide sequence <sup>a</sup>	Effect	. ,	Reference(s) for phosphorylation site
DUT (P33316)	P100S (VAR_022314)	S99 <sup>b</sup> (CDC2)	GPETPAI S <u>P</u> SKRARP	Polymorphism		17081983 8631817
GJA1 (P17302)	P283L (VAR_014101)	S282 <sup>b</sup> (ERK1, ERK2 and MAPK7)	TAPLSPM S <u>P</u> PGYKLV	Polymorphism (rs2228974)		8631994 9535905
PPARG (P37231)	P113Q (VAR_010724)	S112 <sup>c</sup> (ERK2, JNK1 and MAPK8)	AIKVEPA S <u>P</u> PYYSEK	Obesity and polymorphism (rs1800571)	9753710	9030579
RXRA (P19793)	P261L (VAR_014620)	S260 <sup>b</sup> (ERK2 and MAPK7)	NMGLNPS S <u>P</u> NDPVTN	Polymorphism (rs2234960)		12048211

<sup>&</sup>lt;sup>a</sup>The variation sites are underlined and are marked with the bold style.

**Table 3.** Possible examples of type III phosphovariants

Gene name (Swiss-Prot ID)	Variation site (Swiss- Prot variant ID)	Related phos- phorylation site (kinase recognizing it)	Local peptide sequence	Effect	Reference(s) for variation	Reference(s) for phosphorylation site
Panel a: A possible	example of type III phos	phovariant				
PTPN1 (P18031)	P387L (VAR_022014)	S386 (CDC2 and CK2)	LRGAQAA S <u>P</u> AKGEPS	Low glucose tolerance and polymorphism (rs16995309)	15919835	9600099 8491187
Panel b: Possible ex	amples of phosphovariant	s which can be cla	assified as type I or III			
BRCA1 (P38398)	S1217Y (VAR_020695)	S1217	ESSEENL S SEDEELP	Breast cancer and breast-ovarian cancer	14722926	17081983
CASP8 (Q14790)	S219T (VAR_025816)	S219	PREQDSE S QTLDKVY			17525332
CDK2 (P24941)	Y15S (VAR_016157)	Y15	EKIGEGT Y GVVYKAR			1396589 12912980 12972555 15144186
CTNNB1 (P35222)	S33Y (VAR_017619)	S33	QQQSYLD S GIHSGAT	Pilomatrixoma	12027456 10192393	12000790 12114015 11818547
CTNNB1 (P35222)	S37Y (VAR_017627)	S37	YLDSGIH S GATTTAP	Hepatocellular carcinoma	10435629	12000790 12114015 11818547
TEK (Q02763)	Y897S (VAR_008716)	Y897	GACEHRG Y LYLAIEY	Dominantly inherited venous malformations	10369874	11080633
XRCC1 (P18887)	S485Y (VAR_014779)	S485	QDNGAED S GDTEDEL			15066279

The reasons why these variations are classified as type III are detailed in the text.

Protein names which are abbreviated by their gene names: breast cancer type 1 susceptibility proteinBRCA1; caspase 8, CASP8; cyclin dependent kinase 2, CDK2; catenin β-1, CTNNB1; tyrosine-protein phosphatase nonreceptor type 1, PTPN1; angiopoietin-1 receptor (precursorP, TEK; DNA repair protein XRCC1, XRCC1.

<sup>&</sup>lt;sup>b</sup>The removals of the phosphorylation sites by the variation have not been confirmed by experiments. However, the removals of the phosphorylation sites are highly possible because the nearby phosphorylation sites are proved to be recognized by the CMGC group.

<sup>&</sup>lt;sup>c</sup>The removal of the phosphorylation site by the variation has been confirmed by a experiment (12).

If the variations substitute the proline residues at position +1 relative to the phosphorylation sites into other amino acids, the nearby phosphorylation sites recognized by the CMGC kinase group can be eliminated or the efficiency of phosphorylation in that site is significantly decreased. Protein names which are abbreviated by their gene names: deoxyuridine 5'-triphosphate nucleotidohydrolase, mitochondrial (precursor), DUT; gap junction α-1 protein, GJA1; peroxisome proliferator-activated receptor γ, PPARG; retinoic acid receptor RXR-α, RXRA.

72.39%, respectively, for each data set, whereas the sensitivities decreased to 73.24% and 65.76%, respectively (Table 5). At a specificity of >99%, the specificities changed to 95.79% and 96.62%, respectively, and the sensitivities to 23.39% and 20.05%, respectively. Scansite is a widely used web-based prediction software for phosphorylation sites (19). We also tested Scansite with the same data sets. When we applied Scansite with the low-stringency option to the data sets, the

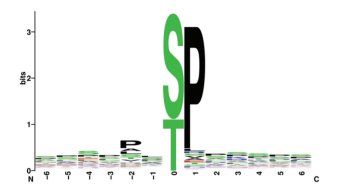


Figure 2. Sequence logos of amino acid sequences near phosphorylation sites recognized by the CMGC kinase group. The horizontal axis represents sequential positions relative to the phosphorylation site. The vertical axis represents decreases in uncertainty. Each letter refers to an amino acid. As the frequency of an amino acid at a given position increases, its height increases.

Table 4. The general performance test: the number of each data set

	Data I <sup>a</sup>		Data	ı II <sup>a</sup>
	(+) <sup>b</sup>	(-) <sup>b</sup>	(+)	(-)
Ser/Thr <sup>b</sup> Tyr <sup>b</sup>	1860 38	926 95	3017 359	315 11

<sup>&</sup>lt;sup>a</sup>Different data sets compiled with mass spectrometer. See the text for the detail explanation for data set I and II.

bThe types of amino acids located at the center of peptides. We anno-

specificities were 52.60% and 57.06%, respectively, and the sensitivities were 84.47% and 83.92%, respectively, whereas with the high-stringency option, the specificities were 96.77% and 95.71%, respectively, and the sensitivities were 16.39% and 13.60%, respectively (Table 6). The performance of PredPhospho with no modification to the specificity is similar to that of Scansite used with the lowstringency option. The performance of PredPhospho with a specificity of >99% was similar with that of Scansite used with the high-stringency option. The data sets were analyzed in the same way with the 18 kinase family models (Tables 5 and 6). The family-wise prediction generally had greater sensitivity and lower specificity than the groupwise prediction.

# Phosphovariants predicted with PredPhospho and Scansite

The numbers of phosphovariants predicted with PredPhospho and Scansite are shown in Table 7. In the Supplementary Data, we present the results for phosphovariants that were predicted with PredPhospho with the >99% specificity option. The sensitivity and specificity of the prediction of type I phosphovariants will be the same as the result shown in Table 5, because we can predict type I phosphovariants simply by the location of phosphorylation sites, with no knowledge of the kinds of kinases involved. The notions of type II and type III phosphovariants include the kinds of kinases that recognize the phosphorylation sites. Therefore, the prediction of type II and III phosphovariants differs from that of type I. Not only the phosphorylation site but also the type of kinase

Table 6. The general performance test of the Scansite for two real data sets

Stringency <sup>a</sup>	Da	ta I	Dat	a II
	Sn (%)	Sp (%)	Sn (%)	Sp (%)
Low	84.47	52.60	83.92	57.06
Medium	48.63	85.21	43.81	87.73
High	16.39	96.77	13.60	95.71

<sup>a</sup>Scansite has three levels of stringency: high, medium and low. High stringency involves low sensitivity and high specificity, whereas low stringency involves high sensitivity and low specificity.

Table 5. The general performance test of the PredPhospho for two real data sets

Specificity <sup>a</sup> Sn		Prediction at	the group level		Specificity		Prediction at t	he family level	
	Da	ta I	Dat	a II		Da	ta I	Data II	
	Sn (%)	Sp (%)	Sn (%)	Sp (%)		Sn (%)	Sp (%)	Sn (%)	Sp (%)
No	79.40	60.62	75.47	61.04	No	95.36	29.29	93.60	30.98
95%	73.24	72.09	65.76	72.39	95%	92.68	42.80	88.95	46.63
97%	53.37	80.22	48.31	79.45	97%	88.46	56.81	83.50	57.36
98%	43.11	89.23	38.92	89.26	98%	82.03	66.80	75.44	62.88
99%	23.39	95.79	20.05	96.62	99%	73.18	72.67	66.73	72.39

<sup>&</sup>lt;sup>a</sup>Options of the specificity. For example, '99%' specificity option mean cutoff value is adjusted for each model to have 99% specificity, and 'No' specificity option means each model has default cutoff value without adjustment of specificity (See supplementary material). Abbreviations: sensitivity, Sn; specificity, Sp.

tated the peptides as (+) if the Ser/Thr or Tyr at the center of the peptides is phosphorylated. On contrary, we designated the peptide as (-) if the center of the peptides is not phosphorylated.

Table 7. The number of the phosphovariants

Specificity	Type I(-)	Type I(+)	Type II(-)	Type II(+)	Type III			
Panel a: P	redicted with	PredPhosph	o at the kina	se group leve	l			
No	1729	2036	5455	4980	5299			
95%	981	1195	1304	1070	986			
97%	613	778	694	542	401			
98%	314	409	329	213	151			
99%	116	150	98	52	21			
Panel b: P	Panel b: Predicted with PredPhospho at the kinase family level							
No	3039	3717	3969	3926	23 955			
95%	2379	2910	2882	2840	8113			
97%	1720	2104	1439	1483	2390			
98%	1268	1551	783	862	1213			
99%	946	1180	539	548	638			
Panel c: Predicted with Scansite								
Low	1581	1852	4255	3773	7697			
Medium	443	498	487	384	152			
High	83	128	35	28	1			

must be identified, because important amino acid residues flanking the phosphorylation sites, which guide kinases to the site, may differ according to the kinase involved. Therefore, the general performance of our prediction of type II and type III phosphovariants will be somewhat different from those shown in Table 5. Instead, the performance of the each kinase-specific prediction can be judged from the performances shown in Supplementary Table S2.

The proportion of phosphovariants predicted by PredPhospho is shown in Table 8. These data were selected with the >99\% specificity option at the kinase family level, and are related to the confirmed phosphorylation sites in human or orthologous proteins, for which kinase information is not yet available. As described in the Supplementary Data, only phosphorylation sites with available kinase information were used as training models for PredPhospho. Hence, the phosphorylation sites in Table 8 are predicted ones with PredPhospho, because the phosphorylation sites were not included in the training data for PredPhospho. If a specific site is shown to be a phosphorylation site in human or orthologous proteins, then the site is definitely located on the surface of the protein, and therefore, is accessible to a kinase. The predicted phosphovariants related to a proven phosphorylation site are more likely to be true than are those related to an unproven phosphorylation site. Numerous phosphorylation sites in humans or other species are constantly being identified. The priority for further research among predicted phosphovariants can be decided based on the confirmation of specific phosphorylation sites.

#### DISCUSSION

Changes in phosphorylation sites cause various diseases by numerous mechanisms. Some proven mechanisms of the phosphovariants shown in Tables 1–3 are related to changes in the protein's affinity for DNA, inducing hyperphosphorylation and the inhibition of ubiquitinization.

For example, microphthalmia-associated transcription factor (MITF) activates the transcription of the tyrosinase gene. The Ser405Pro change in MITF eliminates the phosphorylation site at Ser405 and inhibits the binding of MITF to DNA. As a result, the mutation causes Waardenburg syndrome type IIa, the symptoms of which include depigmentation and sensorineural hearing loss (20). Abnormal phosphorylation can also cause disease by increasing phosphorylation at other sites. The hyperphosphorylation of tau protein induces neurofibrillary tangles and the accumulation of these tangles can result in Alzheimer's disease and frontotemporal dementia (FTD). Paradoxically, serine threonine protein kinase N (PKN) interrupts the phosphorylation of other sites by phosphorylating Ser637 and Ser669 of tau protein (21). The mutations Ser637Phe and Ser669Leu of tau protein eliminate the recognition sites for PKN and induce the hyperphosphorylation of tau protein. FTD and the respiratory failure with dementia are known to be related to Ser637Phe and Ser669Leu, respectively (22,23). Phosphorylation at Ser32 of NF-κB inhibitor α (Swiss-Prot ID, P25963) causes ubiquitinization and results in the activation of NF-kB (24). The Ser32Ile substitution of NF-κB inhibitor α (Swiss-Prot ID, P25963) eliminates the phosphorylation site at Ser32. Consequently, the ubiquitinization of NF- $\kappa$ B inhibitor  $\alpha$  is inhibited by the Ser32Ile variant of NF-κB inhibitor α and NF-κB cannot be activated. The Ser32Ile variant of NF-κB inhibitor α causes autosomal dominant anhydrotic ectodermal dysplasia with immunodeficiency (25). These are a few examples of phosphovariants. Considering the numerous functional roles played by phosphorylation in vivo, there must be many mechanisms by which phosphovariants can cause specific diseases, and these must be identified.

Some phosphovariants do not cause Mendelianinherited diseases, but change an individual's susceptibility to disease. The Pro113Gln substitution (dbSNP id, rs1800571) of peroxisome proliferator-activated receptor gamma (Swiss-Prot ID, P37231), which eliminates the phosphorylation site at Ser112, is known to cause obesity (12). The Pro387Leu substitution (dbSNP ID, rs16995309) of tyrosine-protein phosphatase nonreceptor, type 1 (Swiss-Prot ID, P18031) is associated with type II diabetes mellitus (16,18). We only found these two polymorphic phosphovariants that are related to disease susceptibility. As shown in Tables 1-3, 21 phosphovariants are polymorphisms, and the biological significance of 19 of these polymorphisms is not yet known. In addition, we do not know the biological significance of most of the polymorphic phosphovariants predicted in our study. Considering the importance of phosphorylation in protein function, polymorphic phosphovariants may well be involved in specific diseases or phenotypes.

Apart from the characteristics of the three types of phosphovariants already suggested, there are other fundamental differences between the type I and other phosphovariants. Type I phosphovariants completely add or remove a phosphorylation site because kinases can only donate a phosphor moiety to an amino acid with a hydroxyl group. However, type II and III phosphovariants can significantly affect kinase kinetics without completely

Table 8. Predicted<sup>a</sup> phosphovariants whose phosphorylation sites were confirmed in human or orthologous proteins

Gene name (Swiss-Prot ID)	Site (Swiss-Prot variant ID)	Related phosphorylation site (predicted kinase recognizing it <sup>b</sup> )	Local peptide sequence	Effect	Reference(s) for variation	Reference(s) for phosphoryla- tion site
Panel a: Type I(-) phos	sphovariants					
ACIN1 (Q9UKV3)	S478F (VAR_022033)	S478 (CDK, GSK, and	VQLVGGL S PLSSPSD	Polymorphism		17242355
MECP2 (P51608)	S229L (VAR_018200)	MAPK) S229 (MAPK)	VKMPFQT S PGGKAEG	(rs3751501) Polymorphism	10767337 12872250	(mouse) <sup>c</sup> 17046689 (rat) <sup>c</sup>
PAH (P00439)	S16P (VAR_000869)	S16 (RSK and CAMKL)	PGLGRKL S DFGQETS	Phenylketonuria	1679029 2246858 1301187	7387651 (rat) <sup>c</sup>
Panel b: Type II(-) pho	osphovariants				1301107	
GTSE1 (Q9NYZ3)	R506W (VAR_024154)	S504 (PKC)	PAPQSLL S A $\underline{\mathbf{R}}$ RVSAL	Polymorphism	10591208	16964243
LIG1 (P18858)	P52L (VAR_020194)	S51 (CDK and MAPK)	GVVSESD S <u>P</u> VKRPGR	(rs140054) Polymorphism (rs4987181)		16964243
Panel c: Type III phosp	hovariants					
Gene name (Swiss-Prot ID)	Site (Swiss-Prot variant ID)	Related phosphorylation site (predicted kinase recognizing it)	Local peptide sequence	Removed kinase family <sup>d</sup> (Added kinase family <sup>e</sup> )	Effect (reference(s) for variation <sup>f</sup> )	Reference(s) for phosphorylation site
ABCB11 (O95342)	R698H (VAR_035352)	S701 (AKT, CAMK2, CAMKL, IKK, RSK,	SIRQ <b>R</b> SK S QLSYLVH	AKT, CAMK2, RSK	Polymorphism (16763017)	17242355 (mouse) <sup>c</sup>
ABL1 (P00519)	P810L (VAR_032678)	and PKC) S809 (GSK and MAPK)	MESSPGS S <u>P</u> PNLTPK	MAPK	Polymorphism (17344846)	17081983
AQP2 (P41181)	P262L (VAR_015255)	S261 (IKK and MAPK)	RQSVELH S <u>P</u> QSLPRG	MAPK	Nephrogenic diabetes insipidus (9550615,	16641100 (rat) <sup>c</sup>
CASP8 (Q14790)	S219T (VAR_025816)	S219 (IKK and PIKK)	PREQDSE $\underline{\mathbf{s}}$ QTLDKVY	IKK	15509592) Polymorphism (rs35976359)	17525332
CHGB (P05060)	R178Q (VAR_020287)	S183 (AKT and GRK)	GERGEDS S EEKHLEE	AKT	Polymorphism (rs910122)	16807684
EIF4G3 (O43432)	P496A (VAR_034009)	S495 (MAPK and RSK)	QNLNSRR S <u>P</u> VPAQIA	MAPK	Polymorphism	17081983
MYBPC3 (Q14896)	G278E (VAR_019891)	S275 (CAMK2, PKA,	LSAFRRT S LAGGGRR	(CK2)	(rs35176330) Familial	16964243 9784245
	,	PKC, RSK, and STE20)	_		hypertrophic cardiomyopathy type 4 (12707239)	(chicken) <sup>c</sup>
PKMYT1 (Q99640)	R140C (VAR_019928)	S143 (AKT, CAMKL, CDK, PKC, and RSK)	YAVK <u>R</u> SM S PFRGPKD	AKT	Polymorphism (rs4149796)	17192257
PPP1R12B (O60237)	R836K (VAR_024177)	S839 (CAMK2, IKK, and RSK)	ERLS <u>R</u> LE S GGSNPTT	CAMK2 and RSK	Polymorphism (rs3881953)	17242355 (mouse) <sup>c</sup>
PARK7 (Q99497)	E64D (VAR_020493)	Y67 (Src)	DAKK <u>e</u> gp y dvvvlpg	(PDGFR)	Autosomal recessive early-onset Parkinson disease 7 (15365989	15592455
PNN (Q9H307)	S671G (VAR_023368)	S667 (AKT, GSK, and	LERSHKS S KGG <u>S</u> SRD	GSK	14607841) Polymorphism	17287340
SH3PXD2A (Q5TCZ1)	R1035Q (VAR_030782)	IKK) S1038 (AKT, CAMKL, PIKK, and RSK)	RLAE <u>r</u> aa s QGSDSPL	AKT and RSK	(rs13021, 10095061) Polymorphism (rs3781365)	17525332
WDR91 (A4D1P6)	L257P (VAR_033358)	S256 (IKK and PKC)	RNASLSQ S <u>L</u> RVGFLS	(MAPK)	Polymorphism (rs292592, 15489334 14702039)	16964243

Eight of the type I(-) phosphovariants (VAR 006195, VAR 023368, VAR 023779, VAR 023644, VAR 030238, VAR 020306, VAR 033686, and VAR 027260), and a type III phosphovariant (VAR 020695) were also predicted. However, their detail information are already written in Table 1 and 3.

Protein names which are abbreviated by their gene names: Proto-oncogene tyrosine-protein kinase ABL1, ABL1; ATP-binding cassette sub-family B member 11, ABCB11; Apoptotic chromatin condensation inducer in the nucleus, ACIN1; Aquaporin-2, AQP2; Caspase-8 [Precursor], CASP8; Secretogranin-1 [precursor], CHGB; Eukaryotic translation initiation factor 4 gamma 3, EIF4G3; G2 and S phase-expressed protein 1, GTSE1; Zinc finger protein KIAA1802, KIAA1802; DNA ligase 1, LIG1; Methyl-CpG-binding protein 2, MECP2; Myosin-binding protein C, cardiac-type, MYBPC3; Phenylalanine-4-hydroxylase, PAH; Parkinson disease protein 7, PARK7; Membrane-associated tyrosine- and threonine-specific cdc2inhibitory kinase, PKMYT1; Pinin, PNN; Protein phosphatase 1 regulatory subunit 12B, PPP1R12B; Ribosomal protein S6 kinase alpha-3, RPS6KA3; SH3 and PX domain-containing protein 2A, SH3PXD2A; WD repeat-containing protein 91, WDR91.

<sup>&</sup>lt;sup>a</sup>The prediction was done with the 99% specificity option of PredPhospho at the kinase family level.

<sup>&</sup>lt;sup>b</sup>Kinases that were predicted to recognize the original sequence.

<sup>&</sup>lt;sup>c</sup>The experiment was done in the proteins of other than human. The names of the species are written in the parenthesis.

dRemoved kinases mean that they were predicted not to recognize the variation sequences, while they were predicted to recognize the original sequences.

eAdded kinases mean that they were predicted to recognized the variation sequences, while they were predicted not to recognize the original sequences. The added kinases were written in the parentheses.

<sup>&</sup>lt;sup>f</sup>The reference numbers which are started with 'rs' are dbSNP ID.

changing the kinase's recognition site. For example, the Pro387Leu substitution of PTPN1 removes only 75% of the phosphorylation of Ser386 by CDC2 kinase in vitro, rather than 100% (Table 3) (16).

We can explain phenotypic variations and diseases in terms of phosphovariants in more cases than we have anticipated. Of the human proteins registered in the Swiss-Prot database, 25.5% are phosphoproteins and 60.9% of these phosphoproteins have multiple phosphorylation sites. The protein with the greatest number of confirmed phosphorylation sites is the serine/arginine repetitive matrix protein 2 (Swiss-Prot ID, Q9UQ35), with 195 phosphorylation sites. Although we could only determine 62 phosphovariants with a database search, many more phosphovariants must exist. Furthermore, if we count the mutations that add phosphorylation sites and the type II and III phosphovariants that cannot be found without specific programs, the number of mutations associated with changes in phosphorylation sites is much greater. We did not consider haplotypes in this study, for simplicity. However, if two or more nearby variations are frequently linked, a nearby phosphorylation site can be altered, although each variation individually does not affect the phosphorylation site. Therefore, phosphovariants may account for a much greater proportion of human variation than we have anticipated.

Several issues must be resolved in future studies. First, we regarded nonannotated serine, threonine and tyrosine residues as nonphosphorylated sites. Although we tried not to select false nonphosphorylated sites by using real MS data sets, some of the nonphosphorylated sites might be determined to be phosphorylated sites, in the long run. Second, type II and type III phosphovariants can be interchanged. For example, removed phosphorylation sites associated with type II (-) phosphovariants predicted with PredPhospho may be recognized by kinases that are not included in our prediction models. In such cases, these variations are type III phosphovariants, not type II (-). Conversely, if a phosphorylation site is falsely classified as a site recognized by multiple kinases, instead of by one true kinase, and if a variation affects only some of these kinases, this is a type II phosphovariant incorrectly predicted to be a type III phosphovariant. Other points that must be improved are the low sensitivity achieved with the high-specificity option and the low specificity achieved with the no-specificity option of PredPhospho (Table 5). Moreover, the number of types of kinases that we can predict must be increased and haplotypes should be considered in future studies.

Our method can be used in pathophysiological studies of mutations and in the selection of polymorphisms of clinical and phenotypical importance. Many of the papers that have described the variations, shown in Tables 1–3, did not mention that the variations could be related to changes in phosphorylation sites. This could be attributable to the lack of a specific database that connects mutations with phosphorylation sites, or the lack of a general understanding of the association between phosphorylation and mutation. The type I (+), II and III phosphovariants we have defined cannot be identified simply by database analyses. Specific programs are

required to identify these phosphovariants. Accordingly, many nonsense point mutations whose functional mechanisms are unknown can be reconsidered in terms of phosphovariant. Furthermore, if some mutations are predicted to be phosphovariants with our system, further research will clarify the cause of the associated disease or protein function. Our system can be used to select meaningful variations among endless numbers of newly identified polymorphisms. As sequencing techniques advance, a large number of genetic variations are emerging. At present, comparison of whole genomes of individuals is possible, because the human genome can be sequenced in two months (26). A comparison of phosphovariants between individuals or between species can be undertaken before amino acid variations or nucleic acid variations are compared in whole genomes. A reverse genetic approach for unknown protein functions or phenotypic variations is possible with proven phosphovariants. The screening and prediction of phosphovariants can be a starting point for further research.

#### SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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