# **RESEARCH ARTICLE**

# Linear and Conformational B Cell Epitope Prediction of the HER 2 ECD-Subdomain III by *in silico* Methods

# Manijeh Mahdavi<sup>1</sup>, Hassan Mohabatkar<sup>2\*</sup>, Mehrnaz Keyhanfar<sup>2</sup>, Abbas Jafarian Dehkordi<sup>1</sup>, Mohammad Rabbani<sup>1</sup>

# Abstract

Human epidermal growth factor receptor 2 (HER2) is a member of the epidermal growth factor receptor family of receptor tyrosine kinases that plays important roles in all processes of cell development. Their overexpression is related to many cancers, including examples in the breast, ovaries and stomach. Anticancer therapies targeting the HER2 receptor have shown promise, and monoclonal antibodies against subdomains II and IV of the HER2 extra-cellular domain (ECD), Pertuzumab and Herceptin, are currently used in treatments for some types of breast cancers. Since anti HER2 antibodies targeting distinct epitopes have different biological effects on cancer cells; in this research linear and conformational B cell epitopes of HER2 ECD, subdomain III, were identified by bioinformatics analyses using a combination of linear B cell epitope prediction web servers such as ABCpred, BCPREDs, Bepired, Bcepred and Elliprro. Then, Discotope, CBtope and SUPERFICIAL software tools were employed for conformational B cell epitope prediction. In contrast to previously reported epitopes of HER2 ECD we predicted conformational B cell epitopes P1,: 378-393 (PESFDGDPASNTAPLO) and P2<sub>c</sub>: 500-510 (PEDECVGEGLA) by the integrated strategy and P4: PESFDGD-X-TAPLQ; P5: PESFDGDP X TAPLQ; P6: ESFDGDP X NTAPLQP; P7: PESFDGDP-X-NTAPLQ; P8: ESFDG-XX-TAPLQPEQL and P9: ESFDGDP-X-NTAPLQP by SUPERFICIAL software. These epitopes could be further used as peptide antigens to actively immune mice for development of new monoclonal antibodies and peptide cancer vaccines that target different epitopes or structural domains of HER2 ECD.

Keywords: HER2 receptor-conformational B cell epitope-bioinformatics-cancer therapy-peptide vaccine

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# Introduction

Human epidermal growth factor receptor 2 (HER2), also known as ErbB2, c-erbB2 or HER2/neu, is a 185 kDa protein (p185) involved in signal transduction as well as regulating cell growth. Since the discovery of the role of HER2 in tumorigenesis, it has received great attention in cancer research and vaccine design during the past two decades (Tai et al., 2010). HER2 is a transmembrane glycoprotein contains three distinct regions: N-terminal extracellular domain (ECD), a single transmembrane  $\alpha$ -helix (TM), and an intracellular tyrosine kinase domain. The largest part of HER2, the N-terminal ECD is composed of approximately 600 residues (90–110 kDa) which could be divided into four subdomains (I-IV). Subdomains I (residues 23-217) and III (343-510) created a binding site for the receptor's potential ligands (Lax, 1988). While the cysteine-rich subdomains II (218-342) and IV (511-582) are involved in the homodimerization and heterodimerization (Fiore, 1987; Pietras, 1995). HER2 is highly expressed in a significant proportion of breast, ovarian, gastric, colon, prostate and lung cancers. HER2 overexpression in these cancers often correlates with tumor metastasis, making it an attractive target for cancer immunotherapy (Siyi et al., 2008).

Recently, several monoclonal antibodies (MAbs) against HER2 ECD have been developed for clinical applications of cancer diagnosis and therapy (Yum and Robyn, 2002). Although, the majority of these antibodies can inhibit tumor cell growth with various efficiencies in vitro and in vivo, some of them have not any inhibiting effects or even stimulate tumor cell growth. In addition, the diverse functions of anti HER2 antibodies appear to be correlated to their paratopes specificity (Boyer et al., 1999). It was suggested that anti HER2 antibodies with different epitopes can have distinct effects on HER2 physiological functions and changes its signaling properties (Wang et al., 2004). Pertuzumab, also called Omnitarg is a humanized MAb and HER dimerization inhibitor, binds to the dimerization arm (Subdomain II) of HER2 and prevents its dimerization with other HER family members (Franklin et al., 2004). Trastuzumab also denoted Herceptin, is another recombinant humanized MAb that targets Subdomain IV, close to the transmembrane domain, and in several studies its therapeutic efficacy has been confirmed in HER2 overexpressing metastatic and early stage breast cancer (Cho et

<sup>1</sup>Department of Pharmaceutical Biotechnology, Faculty of Pharmacy, Isfahan University of Medical Sciences, <sup>2</sup>Department of Biotechnology, Faculty of Advanced Sciences and Technologies, University of Isfahan, Isfahan, Iran \*For correspondence: h.mohabatkar@ast.ui.ac.ir

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al., 2003; Park et al., 2008). Hence, these two MAb with clinical effects bind to the HER2 receptor in two different regions corresponding to domain II and IV, which indicate the potential inhibiting effects of other MAbs that target HER2 at different sites (Rockberg et al., 2009). However, more studies in clinical application showed that Patients who have had a significant therapeutic effect by Herceptin treatment started to appear the drug resistant (Calabrich et al., 2008). These results urge researchers to conduct more investigations and develop novel humanized recombinant MAbs for HER2. Afterwards, anti-HER2 MAb muA21 and CH401 were developed (Wang et al., 2000; Ishida et al., 1994). MAb muA21 was found specifically inhibited the growth of the human breast cancer SK-BR-3 cells and MAb CH401 had cytotoxity against HER2 expressing tumor cells. These MAbs were targeted the subdomain I of HER2 ECD.

HER2 is also considered as a target for vaccine development for primary prevention of cancers (Wiwanitkit, 2007). Although passive immunotherapy with trastuzumab is approved for treatment of breast cancer, a number of concerns exist with passive immunotherapy. For instance, the treatment is expensive, has a limited duration of action, and necessitates repeated administrations of the MAb. Active immunotherapy with synthetic peptide B-cell epitopes increases the possibility of generating an adaptive immune response, eliciting protein reactive high affinity MAbs (Garrett et al., 2007). Therefore, there is a need to further explore new epitopes of HER2 ECD and utilize them as peptide vaccine. Identification and selection of suitable epitopes is a time-consuming, expensive and difficult work requiring accurate experimental screening. Hence, cheaper strategies such as computational predictions could be used for faster and cheaper screening. Today, there are many bioinformatics tools available that can be used as primary screening step to identify targets of interest more efficiently (Chen et al., 2011). But there are limitations; it is not possible to develop an exact algorithm, consider to the defective knowledge about the immune response and these methods are working approximately well. Moreover, the prediction results produced by different tools may have distinct differences and multiple tools should be used to gain a consensus result. Therefore, in the current research new linear and conformational B-cell epitopes were predicted by combination of bioinformatics analyses.

B-cell epitopes are defined as regions on the surface of the native antigen that are recognized and bind to B-cell receptors or specific antibodies. B-cell epitopes can be categorized into two types: linear (continuous) epitopes and conformational (discontinuous) epitopes (Zhang et al., 2011). Linear epitopes comprise residues that are continuous in the sequence, while conformational epitopes consist of residues that are distantly separated in the sequence but have spatial proximity. These epitopes are the focus of pathogenesis and immunological research as well as the targets of development of vaccine, cancer therapy and diagnostic reagent (Jiang et al., 2010). Our studies showed that there is no MAb that targets subdomain III of HER2 due to the role of this subdomain in formation of ligand binding site for the receptor; we are trying to 3054 Asian Pacific Journal of Cancer Prevention, Vol 13, 2012

design B cell epitopes in the subdomain III. Therefore, the identification of B-cell epitopes for subdomain III of HER2 ECD can provide important information for the development of new MAb and cancer vaccines that target different epitopes or structural domains of HER2 ECD. In the present study, the linear and conformational B-cell epitopes of HER2 ECD were screened and identified using bioinformatics comprehensive analyses. An integrated strategy was employed for B cell epitope prediction by combining results from sequence and structure based methods and protein-protein interaction tools. Furthermore, we used SUPERFICIAL software that based on protein structures and generates library proposals including linear and nonlinear peptides.

### **Materials and Methods**

#### Prediction of B cell epitopes by integrated strategy

For determination of linear epitopes, the sequence of ECD HER2 receptor was submitted to ABCpred, BCPREDS, Bcepred, Bepipred and Ellipro web servers (Saha and Raghava, 2004: 2006; Larsen et al., 2006; Ponomarenko et al., 2008; Manzalawy et al., 2008a,b). For prediction of discontinuous or conformational epitopes, the PDB structures: 1N8Z, 1S78, 2A91 and sequence based PDB structure obtained by m4t server, a fully automated comparative protein structure modeling server (http://www.fiserlab.org/servers/m4t) was submitted to Discotope 1.2 server (Haste Andersen et al., 2006). Also CBtope was used to predict conformational B cell epitopes using antigen primary sequence (Ansari et al., 2010). Protein-protein interaction sites on the PDB structures were predicted using cons PPISP online tool (Bradford and Westhead, 2005). Solvent accessibility of all the residues was calculated by ASAView (http://www.netasa. org/asaview). Absolute surface area (ASA) of each residue was calculated by DSSP program. These values were transformed to relative values of ASA. Default settings were applied to all the tools used. The regions recognized frequently by four or more than four tools were selected. The protein structures were analyzed and viewed with PyMOL V1.

#### Secondary Structure Prediction

The secondary structure was predicted using PHD (http://npsa-pbil.ibcp.fr/cgi-bin/npsa\_automat.pl?page=/NPSA/npsa\_phd.html) and PSIPRED (http://bioinf.cs.ucl. ac.uk/psipred/) fold servers. The secondary structure prediction results were used for conformational B-cell epitope identification by integrated strategy.

### Theoretical physico-chemical properties of the peptides

The theoretical physicochemical properties of the synthetic peptides (such as the ionic status, calculated as the isoelectric point, and the hydrophobicity, measured as the grand average of hydropathicity (GRAVY) index) were analyzed using the Prot Param algorithm (http://www.expasy.ch/tools/protparam. html). The GRAVY index indicated the hydrophobicity of the peptide and was calculated as the sum of the hydropathy values (Kyte and Doolittle parameters) of the composing amino

acids, divided by the number of residues in the sequence. Peptides with positive GRAVY index are hydrophilic whereas peptides with negative GRAVY index are hydrophobic (Lebreton et al., 2011).

# Prediction of B cell epitopes by SUPERFICIAL software

We used SUPERFICIAL (http://bioinformatics.charite. de/superficial) a structure-based program for designing of peptide libraries that mimic the whole surface or a particular region of a protein. It automatically defines the protein surface, using preset default values useful to a range of proteins. SUPERFICIAL uses protein structures (PDB codes) as input and generates library proposals consisting of linear and nonlinear peptides. Neighboring sequence segments are linked by short spacers to conserve local conformation. All generated peptides are listed within a table. Such a structure-based peptide library provides the source for chemically-prepared peptides to identify and characterize binding sites, respectively (Goede et al., 2005).

## Sequence Alignment

Amino acid sequence of ECD of human and mouse ErbB2 receptor and peptides (P1<sub>c</sub>, P2<sub>c</sub>) was aligned using Cobalt Constraint-based Multiple Protein Alignment Tool (http://www.ncbi.nlm.nih.gov/tools/cobalt/cobalt. cgi?link\_loc=BlastHomeAd) and The Universal Protein Resource (UniProt)/Align (http://www.uniprot.org/) with default parameters. The output was manually checked to optimize the alignment.

# Results

## Prediction of B cell epitopes by integrated strategy

The predicted linear B-cell epitopes of subdomain III of HER2 ECD are shown in Table 1. For each peptide, sequence, position, molecular weight, isoelecteric pH and gravy score was indicated. As shown in Table 1, P5 and P15 are recognized frequently by four and five tools used in this work. These peptides are related to regions 379-394 and 491-506, respectively. P8 and P12 were only identified by ABCpred.

Furthermore, in CBtope software four conformational B-cell epitope regions were identified which two of them were in the region of linear B-cell epitopes (376-394 and 492-508). Also conformational B-cell epitope (376-394) was identified by discotope server (Table 2). Further analyses for protein-protein interaction sites and solvent accessibility of all the residues on the PDB structures using cons PPISP and ASAView online tools defined that B cell epitopes of 378-393 (PESFDGDPASNTAPLQ), P1<sub>c</sub> and 500-510 (PEDECVGEGLA), P2<sub>c</sub> have higher solvent accessible and their residues are exposed on the surface (Figure 1). These are important factors for immunogenicity of an antigen or synthetic peptide. Around 43% of the 170 residues (in subdomain III) were non polar, 22% were charged residues and 34% were recorded as polar residues for subdomain III of HER2 ECD. The relative surface accessibility values calculated by ASAView showed 34.91% residues with greater than 30% whereas 27.81% residues were with greater than 50% solvent accessibility. Cons PPISP online tool was determined buried residues, 40.24% of residues in subdomain III of HER2 ECD were





 Table 1. The 16mer Peptides Linear B-cell Epitopes Predicted from Subdomain III of HER2 ECD by Different Linear B-Cell Epitope Prediction

Gravy score	pI	Mw (Da)	Softwares <sup>b</sup>	Position	Sequence	Peptide <sup>a</sup>
-1.337	4.87	1783.9	1,2,3	321-336	EVTAEDGTQRCEKCSK	*P1
-0.194	6.76	1818	1,3	343-358	YGLGMEHLREVRAVTS	*P2
0.231	8.24	1679.9	1,4,5	355-370	AVTSANIQEFAGCKKI	P3
0.4	6.06	1802.1	1,5	367-382	CKKIFGSLAFLPESFD	P4
-0.925	3.49	1645.7	1,,2,3,5	379-394	ESFDGDPASNTAPLQP	*P5
-0.35	4	1739.9	1,2,4	385-400	PASNTAPLQPEQLQVF	*P6
-0.188	3.56	1830	3,5	392-407	AWPDSLPDLSVFQNLQ	*P7
0.006	3.57	1857	<sub>1</sub> 10	<b>0.0</b> <sub>403-418</sub>	LEEITGY <del>LYISA</del> WPDS	P8
0.163	3.56	1838	1,2	411-426 6.3	<b>YASAWPDSLPDLSVFQ</b>	P9
0.269	10.84	1810.1	1,5	428-443	LQVIRGRILHNGAYSL	P10
0.406	9.6	1757	1,5	448-463	LGISWLGLRSLRELGS	P11
0.713	7.01	1733	1 /	5.0 <sub>461-476</sub>	LGSGLALIHHNTHLCF.	P12
0.35	7.09	1954.2	1,3,4	467-482	LIHHNTHLCFVHTVPW	P13
-0.806	6.89	1945.1	1,2,5	477-492 56.3	VH18VPWDQLFRNPHQA	*P14
-0.625	4.65	1752.9	1,2,3,4,5 <sub>E</sub>	a a <sup>491-506</sup>	QALLHTANRBEDECVG	*P15
-0.312	4.79	1975.2	2,3,5	519-535	VEECRVLQGLPREYV3143	*P16





30.0

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#### **Table 2. Predicted Conformational B Cell Epitopes**

Sequence <sup>a</sup>		Residue no	Position	Softwares			
FLPESFDG	DPASNTAPLQI	P 17	376-394	Cbtope, Discotope			
FQNLQVI	RGRILHN	9	425-438	CBtope			
ALLHTAN	RPEDECVGEG	14	492-508	CBtope			
GHCWGPC	GPTQCVNCSQF	LRGQE17	518-539	CBtope			
Protein-protein interaction sites and solvent accessibility of all the residues (cons PPISP and ASAView online tool)							
Gravy score pI Mw (Da) Position Residue no Sequence <sup>a</sup> Peptide <sup>b</sup>							
-1.773	3.49 1140.1 378	8-393 11 P	ESFDGDI	PASNTAPLQ *P1			

3.5 1015 500-510 10 PEDECVGEGLA -0.66 \*P2

<sup>a</sup>B cell epitope residues are indicated as Bold and underlined, <sup>b</sup>Hydrophobic Peptides are stared (i. e \*P1)



Figure 2. Screenshot of SUPERFICIAL Showing the 3D View of Conformational B Cell Epitope PESFDGD-X -TAPLO (378-385, 389-393) on Subdomain III of Her 2 ECD. The epitopes are depicted in green color. b and c) the 100.6 pitopes was obtained by integrated strategy (Table 1). 3D view of this epitope on the PDB structure 1N8z by Pymol software. The epitopes are depicted in pink color. In b the epitope is shown in HER2 ECD structure and in c the epitope is magnified on subdomain III.

buried and the others (61.24%) had no contact together. Secondary Structure Prediction

Secondary structure prediction with PHD server identified 10 a helices (12.73%), 72.56 % random coil, 14.71% extended strand and no  $\beta$  sheet in HER2 regions recognized by PSIPRED from amino acid sequence 339-341 (IQE) and 372-376 (PEQLQ) were related to subdomain III of HER2 ECD. Furthermore on PHD server, residues 336-342 (SANIQEF), 399-409 (DLSVFQNLQVI), 372-385 (PEQLQVFETLEEIT) and 433-439 (LRSLREL) were recognized in alpha helical region of subdomain III. The results of PSIPRED, 339-341 (IQE) and 372-376 (PEQLQ) is similar to the results of PHD server, 336-342 (SANIQEF), 372-385 (PEQLQVFETLEEIT).

Prediction of B cell epitopes by SUPERFICIAL software The PDB structures 1N8Z, 1S78, 2A91 and sequence based PDB structure obtained by m4t server, was submitted to SUPERFICIAL software. This program uses protein structures as input and generates library proposals consisting of linear and nonlinear peptides (Table 3 and

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### Table 3. The Linear B-Cell Epitopes Predicted from Subdomain III of Her 2 ECD by SUPERFICIAL Software.

PDB code	Position	Res	idue no	Sequence	Pep	tide
1N8Z, 1S78, 2A91, M4T	342-355	14	CYGLO	GMEHLREV	'RA	P1
1S78, 1N8Z, 2A91, M4T	362-369	8	QEFAG	СКК		P2
1S78, 2A91, M4T	382-390	9	DGDPA	SNTA		P3
1N8Z, M4T	394-401	8	PEQLQ	VFE		P4
1S78	434-439	6	RILHN	G		P5
1S78, 1N8Z, 2A91, M4T	483-489	7	DQLFR	NP		P6
1S78, 1N8Z, 2A91, M4T	495-518	24	HTANF	RPEDECVG	EG	
			LACHO	QLCARG		P7
1S78	529-533	5	VNCSC	2		P8



Figure 3. Multiple alignments between peptides (P1<sub>c</sub>, P2<sub>c</sub>) and ErbB2 ECD of human and mouse using **Cobalt Constraint-based Multiple Protein Alignment** Tool.

4).

In Table 3 and 4, linear and nonlinear peptides were shown, respectively. Linear peptides were predicted by SUPERFICIAL software was similar to Linear B cell In Table 4, conformational B-cell epitopes P4-P9 were in the region  $6.31 \text{ P}_{C}$ :  $3498-394 \text{ of }2013 \text{ R} 2 \text{ E}_{CD}$  subdomain III and also P13-15 was nearly similar to P2. SUPERFICIAL 75. Predicted new conformational B cell epi250 or nonlinear peptides such as P1-3 and P10-12. To preserve their conform**56** and the gaps between the peptide-fragments 50. Ore filled with linkers, short anging acid sequences derived from the LIP (Loops in Proteins) database (Michalsky et al., 2003). Certainly sometimes amino acids were represented by the character "X" that could be replaced ECD whereas PSIPRED server predicted 3  $\alpha$  helices, 25.9 praxis by poly-Alarine and/or Glycine instead of 24 extended strand and no  $\beta$  sheets. The alpha helical using the **31.3** ker. Figure 2 is showing the **31.3** view of the using the **31.3** ker. Figure 2 is showing the **31.3** view of the predicted conformational B-cell epitope PESFDGD-X 6TAPLQ (378-385, 389-393) in SUPERFICIA and Pymol softwares. emissior



The sequence alignment between ECD of human and mouse ErbB2 receptor is form using UniProt was shown 85 \$2% identity, 539 identical positions and 63 similar positions. Additionally the sequence homology of predicted HER2 peptides between the human and mouse ExbB2 sequences (Figure 3). It was shown these peptides and homology of 100% with HER2 ECD. This alignment was also indicated 64 and 50% homology of peptides  $P1_c$  and  $P2_c$  with mouse ErbB2 ECD, respectively. These results were described the rate of foreignness of HER2 ECD for immunization of mouse. It should be noted the foreignness is a very important factor

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PDB code	Position	Sequence	Peptide
1S78B, 1N8Z, m4t	362-369, 400-404	QEFAGCKK-XXXX-FETLE	P1
2A91	362-368, 396-400	QEFAGCK-XXXXX-QLQVF	P2
2A91	362-369, 399-404	QEFAGCKK-XXXX-VFETLE	P3
1S78A, 1N8Z	378-384, 389-393	PESFDGD-X-TAPLQ	P4
1S78A	378-385, 389-393	PESFDGDP X TAPLQ	P6
1S78A	379-385, 388-394	ESFDGDP X NTAPLQP	P6
2A91	378-385, 388-393	PESFDGDP-X-NTAPLQ	P7
2A91	379-383, 389-396	100.Q <sub>SFDG-XX-TAPLQPEQL</sub>	P8
2A91	379-386, 388-394	ESFDGDI6.3-NTAPLOP	Р9
1S78B, 1N8Z, m4t	395-404, 364-368	EQLQVFETLE -XXXX-FAGCK	P10
1N8Z, m4t	396-400, 362-368	QLQVF-XXXX-QEFAGCK	э <b>го</b> Р11
1S78B, 1N8Z, m4t, 2A91	397-404, 364-369	75.UQVFETLE-XXXX-FAGCKK	25.0 P12
1N8Z, 2A91	509-518, 523-528	LACHQLCARG-XXX-PGPTQC	P13
1N8Z	510-515, 523-529	ACHQL (5% XX-PGP1QCV	P14
1N8Z, 2A91	510-518, 523-530	50.0 CHQLCARG-XXX-PGPTQC542	P15
			31.3

Table 4. The Conformational or Non-Linear B-cell Epitope Regions Predicted from Subdomain III of Her 2ECD by SUPERFICIAL Software

for immunogenicity of an antigen or synthetic peptide (Richard et al., 2003).

# Discussion

Our study is the first to report the identification of novel conformational B-cell epitopes of HER2 ECD subdomain III. Despite of majority of epitopes being conformational, most of the computational methods and databases centered at the sequential epitopes (Vita et al., 2010). Linear epitope prediction methods can be categorized into physicochemical property, Hidden Markov Mode (HMM) and Artificial Neural Networks (ANN) based (Saha and Raghava, 2004; Larsen et al., 2006; Saha and Raghava, 2006). For this purpose, ABCpred, BCPREDs, Bepired, Bcepred and Ellipro were used in this research. If the tertiary structure of an antigen is known, there are improved methods for the prediction of conformational B cell epitopes. Examples are SUPERFICIAL software and Discotope web server (Goede et al., 2005; Haste Andersen et al., 2006; Moreau et al., 2008; Ponomarenko et al., 2008). These are based on features like flexibility, solvent accessibility and amino acid propensity scales (Kulkarni-Kale et al., 2005; Haste et al., 2006).

In the current study attempts have been made to predict linear and conformational B cell epitopes in HER2 receptor from both primary sequence and tertiary structure. Two different methods were used based on bioinformatics analyses. In the first method, integrated strategy, the best peptides were the linear B-cell epitopes P5: 379-394 and P15:491-506 (Table 1) and conformational B-cell epitope P1<sub>c</sub>: 378-393 and P2<sub>c</sub>: 500-510 (Table 2) which was different from the previously reported epitopes for the subdomain III of the receptor. Both of the linear and conformational epitopes were approximately in the identical regions of HER2 ECD. Based on secondary structure prediction results (PSIPRED), position 378-393 of HER2 ECD had an alpha helical structure. In the second method, SUPERFICIAL software were predicted many new conformational peptide linked by the linkers (Table 4) and also this software is predicted conformational B-cell epitope  $P1_c$ : 378-393 and  $P2_c$ : 500-510 in the form of two

segments or linear B-cell epitopes that are linked by the **25.0** inker (Figure 2 and Table 3).

30.0

30.0

30.0

None

In the majority 38 the past studies on HER2 receptor, either Linear B cell epitopezagr T-cell epitopes were used as synthetic peptide or antigen, while the vaccine eptides were designed to be chimeric with multi-epitopes of B-cells and helpet T cells (Pakappaga i et al., 2000; Montgomgry et al., 2005). Synthetic periodes can be presented to in different ways such as zonjugation to different carrier protains, incorporation into liposomes, covalent ginjugation to fatty gicids and generation of multiple antigen per kides (M P) (Haro and Gomara, 2004; Vejeet al., 20 E). Miyak et al. (2011) generated 22 kinds 🕏 synthet 😇 MAPs containing 20 amino acid residues based on the region covering from N: 143 to N: 370 of HE 2 ECD. They identified a new epitope of MAb CH401 against HER2 extracellular domain (N: 167-175), and evaluated the effect of active immunization of the 20mer peptide containing the epitope (CH401 peptide). The 20mer peptide (N: 163-182) including CH401 epitope (CH401 epitope peptide) was immunogenic, causing a high titer of peptide-specific IgG antibody in immunized BALB/c mice because it contained epitopes for both B-cells and helper T-cells. In contrast to the present study, these epitopes were located on subdomain I of HER2 ECD. In addition, Dakappagari et al. (2003) identified two linear HER2 ECD B cell epitopes (316-339 and 485–503) based on computer aid analyses. These epitopes incorporated to the measles virus fusion (MVF) 'promiscuous' T cell epitope via a four-residue linker sequence. They induced high titrated Abs inhibiting the growth of human breast cancer cells (Dakappagari et al., 2003). Moreover, Garrett et al. (2007) reported phase I clinical trial using the first generation peptide vaccines MVF 316-339 and MVF 628-647. They also used peptide constructs MVF 613-626, MVF 563-598 and MVF 597-626. All epitopes were immunogenic in FVB/n mice and 597-626 epitope significantly reduced cancer growth in transgenic BALB-neuT mice. Besides, Wiwanitiki (2007) predicted B cell epitopes of HER2 oncoprotein by ABCpred prediction server. He determined the peptide 947SRMARDPQRFVVIQNE963 with the

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best binding affinity. Thus, the finding of present work is different from those predicted epitopes or peptides of the previous studies (Dakappagari et al., 2003; Garrett et al., 2007; Wiwanitiki, 2007; Furthermore et al., 2009; Miyako et al., 2011) generated antibodies to linear epitopes of the HER2 that effect cell growth by a strategy based on polyclonal antibodies. The functional antibodies with growth inhibition properties located on domain II and III of the receptor, respectively. One of the epitopes mapped to domain II overlapped with the conformational epitope recognized by pertuzumab (266-296), while similar to our work, another epitopes on the surface of the domain III were close to each other in a linear stretch of 25 amino acids LPESFDGDPASNTAPLQPEQLQVF. Jasiinska et al. (2003) also identified seven B cell epitopes on sequence of HER2 ECD by computer analyses. The peptides (P1: 115-132AVLDNGDPLNNTTPVTGA; P2: 149-162LKGGVLIQRNPQLC; P3: 274-295 Y N T D T F E S M P N P E G R Y T F G A S; P4: 378-398PESFDGDPASNTAPLQPEQLQ; P5: 489-504PHQALLHTANRPEDE; P6: 544-560CRVLQGLPREYVNARHC; P7: 610-623YMPIWKFPDEEGAC were used for immunization, successfully induced humoral immune response with anti-tumor activity in an animal model. The linear peptide P5: 379-394 that we are predicted is similar to epitope P4: 378-398 which is used in Jasinska et al. (2003) research. In another study, Dakappagari et al. (2005), designed a complex immunogen and conformational B-cell epitope peptide vaccine derived from the extracellular domain of HER2-(626-649) that represents a three-dimensional epitope. They successfully introduced two disulfide bonds into this sequence. Their study demonstrate the feasibility and importance of designing conformational epitopes that mimic the tertiary structure of the native protein for eliciting biologically relevant anti-tumor antibodies. Such approaches are a preequipment to the design of effective peptide vaccines and immunogenic synthetic peptides. In the present work, new conformational B cell epitopes P1<sub>c</sub>: 378-393 (PESFDGDPASNTAPLQ) and P2<sub>c</sub>: 500-510 PEDECVGEGLA (Table 2) were predicted by bioinformatics analyses (integrated strategy and SUPERFICIAL software). These peptides are discontinuous parts of region 378-393 and 500-510 of HER2 ECD which were predicted with Linear B cell epitope tools.

Tolerance to self antigens such as HER2, may limit a functional immune response to whole protein based antigens due to activation of suppressor T cells that maintain tolerance to host Antigens (Disis. and Cheever, 1997; Sakaguchi, 2000; Bonilla and Oettgen, 2010), while immunizing with peptides derived from self antigens may be an effective means of circumventing tolerance (Ercolini et al., 2003). However, it should be noted that peptides with a sequence identical to that of self antigen will not be able to break the threshold of self tolerance status and specific lymphocytes is expected to be deleted during the establishment of self identity. Mittelman et al. (2002) analyzed the similarity level of HER2 peptides to the host's proteome and understand the relationship between molecular mimicry and peptide immunogenicity. They reported low level of sequence similarity to the host's proteome plays an important role in shaping the pool of B cell epitopes and determining peptide immunogenicity. This finding has important implications in HER2 cancer therapy. For this purpose, in our study the foreignness or similarity level of peptides were evaluated by multiple sequence alignment. The results indicated lower similarity of peptides with mouse (Figure 3) compared to HER2 ECD. Furthermore, the peptides, P1<sub>c</sub> and P2<sub>c</sub>, are indicated high degree of foreignness (50 and 36%, respectively) for immunization of mice.

In conclusion, the findings of present work using bioinformatics analyses could be applied in MAb production, cancer therapy, developing vaccine design and diagnostic tools reducing the time and minimizing the total number of required tests to find possible proper epitopes. In the next step, in vitro synthesis of determined peptides and in vivo experimental studies are essential for assurance of the predicted epitopes. In these experiments, identification of epitopes on HER2 that are either stimulatory or inhibitory is very important for development of strategies to better manipulate the Ab response for therapeutic benefit. Thus, selection of suitable epitopes from specific regions of HER2 capable of inducing cancer inhibitory MAbs and careful elimination of epitopes that stimulate cancer cell growth, will be identified candidate peptides that could provide beneficial effects.

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