pGAPZ A, B, and C pGAPZ α A, B, and C

Version F September 3, 2002 25-0174

pGAPZ A, B, and C pGAPZ α A, B, and C

Pichia expression vectors for constitutive expression and purification of recombinant proteins

Catalog nos. V200-20 and V205-20



www.invitrogen.com tech_service@invitrogen.com

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Technical Services

Invitrogen provides Technical Services to all of our registered *Pichia* Expression Kit users. Please contact us if you need assistance with the *Pichia* Expression Kit.

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Important Information

Contents

This manual is included with the following catalog numbers:

	Catalog no.	Item	Contents			
	V200-20	pGAPZ A, B, and C	20 µg each vector, lyophilized			
		· · ·	X-33/pGAPZ B/lacZ Pichia strain			
			expressing β -galactosidase			
	V205-20	pGAPZ α A, B, and C	20 µg each vector, lyophilized			
		• • •				
Shipping/Storage	The lyophilized ve	ectors are shipped at roor	n temperature and stored at -20°C.			
Materials Supplied by the User	For the procedures described in this manual, you will need the following reagents and equipment. Additional reagents may be required.					
,	Equipment					
	• Pichia Expressi	on Manual				
	Microbiological	l equipment				
	 Electroporation 	device and 0.2 cm cuvet	tes or reagents for LiCl transformation			
	• 16°C, 37°C, and	d 65°C water baths or ten	perature blocks			
	• 30°C and 37°C	shaking and non-shaking	incubators			
	 Hemacytometer Microtiter plate 	s (optional)				
	Reagents	s (optional)				
	• Pichia host strain (X-33, GS115, KM71, or SMD1168)					
	 Electrocompetent or chemically competent <i>E. coli</i> (must be <i>rec</i>A, <i>end</i>A) for transformation 					
	Restriction enzymes and appropriate buffers					
	Sequencing primers to confirm proper insertion of your gene of interest					
	Agarose and low-melt agaroseGlass milk					
	• Sterile water					
	CIAP (calf intes Dhanal/ablarafa	stinal alkaline phosphatas	e, I unit/µI), IOX CIAP Buffer			
	 3 M sodium ace 	niii tate				
	• 100% ethanol 8	30% ethanol				
	• T4 Ligase (2.5)	units/ul), 10X Ligation B	uffer (with ATP)			
	Low Salt LB me	edium (see page 18 for re	cipe)			
	• Zeocin [™] antibio	otic (see page 5 for order	ng information)			
	• Low Salt LB pla	ates containing 25 µg/ml	Zeocin [™]			
	• YPD medium					
	• YPDS plates co	ntaining 100 µg/ml Zeoc	in [™] plates (see page 30 for recipe)			
	• 50 ml conical centrifuge tubes and 15 ml polypropylene tubes					
	• YNB medium c	ontaining 0.5% glucose (see page 30 for recipe)			
	• ProBond [™] Puri	tication System (Catalog	no. K850-01, optional)			

Important Information, continued



Pichia users should already have a *Pichia* Expression System manual. Procedures for analysis of recombinants are described in the *Pichia* manual. For more information, please call Technical Service (page 34).

Other Pichia Products

Other Pichia products available from Invitrogen are described below:

Item	Purpose	Quantity	Catalog no.
Pichia Expression Kit	Original complete kit for gene expression in <i>P. pastoris</i>	1 kit	K1710-01
EasyComp™Kit	Rapid preparation and transformation of competent <i>P. pastoris</i> cells	1 kit	K1730-01
EasySelect [™] <i>Pichia</i> Expression Kit	Complete kit for easy selection of recombinants on Zeocin ^{TM} and gene expression in <i>P. pastoris</i>	1 kit	K1740-01
Multi-Copy <i>Pichia</i> Expression Kit	Complete kit for multiple copy gene insertion into <i>P. pastoris</i>	1 kit	K1750-01
pPICZ A, B, and C	For inducible expression on methanol, selection on Zeocin [™] , and purification of intracellular proteins	20 µg each	V190-20
pPICZα A, B, and C	For inducible expression on methanol, selection on Zeocin [™] , and purification of secreted proteins	20 µg each	V195-20

Introduction

Overview			
Introduction	The glycera expressed at the gene (<i>G</i> , to express re carbon sour promoter (P	Idehyde-3-phosphate t high levels in many (AP) encoding the GA ecombinant proteins t ce used (Waterham <i>et</i> (GAP) can be slightly h	dehydrogenase (GAPDH) enzyme is constitutively organisms, including <i>Pichia pastoris</i> . The promoter of PDH protein has recently been characterized and shown o high levels in <i>Pichia pastoris</i> , depending on the <i>al.</i> , 1997). The level of expression seen with the <i>GAP</i> igher than that obtained with the <i>AOX1</i> promoter.
	The pGAPZ the <i>GAP</i> pro Proteins car for detection ProBond [™]). encoding th supplied in and/or the N selectable m	A, B, and C vectors omoter to constitutive a be expressed as fusion and a polyhistidine to In addition, pGAPZo e Saccharomyces cere three reading frames to I-terminal secretion sin marker, Zeocin [™] , which	(2.9 kb) and pGAPZ α A, B, and C (3.1 kb) vectors use ly express recombinant proteins in <i>Pichia pastoris</i> . ons to a C-terminal peptide containing the <i>myc</i> epitope tag for purification on metal-chelating resin (i.e. α produces proteins fused to an N-terminal peptide <i>evisiae</i> α -factor secretion signal. Both vectors are o facilitate in frame cloning with the C-terminal tag gnal. Selection of these vectors is based on the dominant h is bifunctional in both <i>Pichia</i> and <i>E. coli</i> .
Pichia Strains	The followi	ng <i>Pichia pastoris</i> str	ains can be used with the pGAPZ and pGAPZ α vectors:
	<i>Pichia</i> Strain	Genotype	Purpose
	X-33	wild type	Expression of recombinant proteins from vectors with Zeocin [™] resistance as the only selectable marker (i.e. pGAPZ and pGAPZα).

GS115

KM71

SMD1168

his4

his4, aox1::ARG4

his4, pepA

continued on next page

Expression of recombinant proteins from vectors containing the *HIS4* or Zeocin[™] resistance gene as

Expression of recombinant proteins from vectors

containing the *HIS4* or Zeocin[™] resistance gene as the selectable marker(s) in a Mut^S background when using methanol induction for expression. Protease deficient strain for the expression of

recombinant proteins from vectors containing the HIS4 or ZeocinTM resistance gene as the selectable

the selectable marker(s).

marker(s).

Overview, continued

Experimental Process

The following table describes the overall experimental process.

Step	Action	Page
1	Propagate pGAPZ A, B, and C and/or pGAPZ α A, B, and C by transformation into a <i>rec</i> A, <i>end</i> A1 <i>E. coli</i> strain such as TOP10F', DH5 α , or JM109.	10
2	Develop a cloning strategy to ligate your gene into one of the pGAPZ α or pGAPZ vectors in frame with the α -factor secretion signal and/or the C-terminal tag.	9-17
3	Transform into <i>E. coli</i> and select transformants on low salt LB plates containing 25 μ g/ml Zeocin TM .	18
4	Analyze 10-20 transformants by restriction mapping or sequencing to confirm in frame fusion of your gene with the α -factor secretion signal and/or the C-terminal tag.	19
5	Purify and linearize the recombinant plasmid for transformation into <i>Pichia pastoris</i> .	20
6	Transform your <i>Pichia</i> strain and plate onto YPDS plates containing 100 µg/ml Zeocin [™] .	21
7	Select Zeocin [™] -resistant transformants.	22
8	Test for expression of the gene of interest.	23
9	Optimize expression of your gene.	25
10	Purify your fusion protein on metal-chelating resin (i.e. ProBond [™]).	26

ProBond[™] Resin

We recommend that you use the ProBond[™] Purification System (Catalog no. K850-01) for purifying fusion proteins expressed using the pGAPZ or pGAPZα vectors. **Please note that instructions for equilibration of and chromatography on ProBond[™] resin are contained in the ProBond[™] Purification System Kit.**

If you are using a metal-chelating resin other than ProBond[™], please follow the manufacturer's recommendations for fusion proteins expressed in yeast.

Recombination and Integration in Pichia

Introduction Linear DNA can generate stable transformants of Pichia pastoris via homologous recombination between the transforming DNA and regions of homology within the genome (Cregg et al., 1985; Cregg et al., 1989). Such integrants show extreme stability in the absence of selective pressure even when present as multiple copies. Note that single crossover events (insertions) are much more likely to happen than double crossover events (replacements). Multiple insertion events occur spontaneously at about 1-10% of the single insertion events. Gene Insertion at Gene insertion events at the GAP promoter locus arise from a single crossover event between the locus and the P_{GAP} region on the pGAPZ or pGAPZ α vectors. This results in the GAP Promoter the insertion of one or more copies of the vector upstream or downstream of the GAP locus. The figure below shows the result of an insertion of the plasmid 5' to the intact GAP promoter locus and the gain of P_{GAP}, your gene of interest, and Zeocin[™] resistance. This also occurs with non-linearized plasmid and plasmid that religates, although at a lower frequency. Pichia Genome (HIS4 or his4) GAF Zeo **Expression Cassette**

Recombination and Integration in Pichia, continued

Multiple Gene Insertion Events

Multiple gene insertion events at a single locus in a cell do occur spontaneously with a low, but detectable frequency--between 1 and 10% of all selected Zeo^R transformants. Because of the low frequency of multiple gene insertion events, you will need to screen hundreds to thousands of Zeocin[™]-resistant transformants to locate these "jack-pot" clones. We recommend that you use electroporation to generate Zeo^R transformants for screening.

Please note that multiple gene insertion with the pGAPZ and pGAPZ α has not been functionally tested by Invitrogen. However, since the backbone of these vectors are identical to pPICZ and pPICZ α , a multiple copy event can occur at the *GAP* promoter region locus. Multiple gene insertion events can be detected by quantitative dot blot analysis, Southern blot analysis, and differential hybridization. Please see a *Pichia* Expression Manual for a protocol to screen for multiple inserts.



Zeocin™

Zeocin™

ZeocinTM and the *Sh ble* gene, which encodes the ZeocinTM resistance factor, are used for selection in mammalian cells (Mulsant *et al.*, 1988)plants (Perez *et al.*, 1989); yeast (Baron *et al.*, 1992; Wenzel *et al.*, 1992); and prokaryotes (Drocourt *et al.*, 1990). Suggested concentrations of ZeocinTM for selection *E. coli* and *Pichia pastoris* are listed below:

Organism	Zeocin [™] Concentration and Selective Medium
E. coli	25 µg/ml in Low Salt LB medium*
Pichia pastoris	100 µg/ml in YPDS medium

*Efficient selection requires that the concentration of NaCl be no more than 5 g/liter (< 90 mM).

For more information on Zeocin[™] and Zeocin[™] resistance, please see page 31.



Any *E. coli* strain that contains the complete Tn5 transposable element (i.e. DH5 α F'IQ, SURE, SURE2) encodes the *ble* (bleomycin) resistance gene. These strains will confer resistance to ZeocinTM. For the most efficient selection it is highly recommended that you choose an *E. coli* strain that does not contain the Tn5 gene (i.e. TOP10F', DH5, DH10, etc.).

Purchase of Zeocin[™] ZeocinTM can be purchased from Invitrogen in 1 gram (Catalog no. R250-01) and 5 gram (Catalog no. R250-05) quantities. For your convenience, the drug is prepared in autoclaved, deionized water and aliquoted into 1.25 ml aliquots at a concentration of 100 mg/ml. The stability of ZeocinTM is guaranteed for six months, if stored at -20°C.

Vectors

Features

The vectors pGAPZ A, B, and C and pGAPZ α A, B, and C contain the following elements. All features have been functionally tested.

Feature	Benefit
GAP promoter	Allows constitutive, high-level expression in <i>Pichia</i> (Waterham <i>et al.</i> , 1997)
	Targets plasmid integration to the GAP locus
α-Factor Secretion Signal (pGAPZα A, B, and C only)	Encodes the native <i>Saccharomyces cerevisiae</i> α -factor secretion signal that allows for efficient secretion of most proteins from <i>Pichia</i> (Cregg <i>et al.</i> , 1993)
Multiple cloning site with unique restriction sites	Allows insertion of your gene into the expression vector
C-terminal <i>myc</i> epitope (Glu-Gln-Lys-Leu-Ile-Ser-Glu- Glu-Asp-Leu)	Permits detection of the fusion protein by the Anti- myc Antibody (Catalog no. R950-25) (Evans <i>et al.</i> , 1985)
C-terminal polyhistidine tag	Encodes six histidine residues that form a metal- binding site for affinity purification of recombinant protein (i.e. with ProBond [™] , Catalog no. K850-01) and detection by the Anti-His(C-term) Antibody (Catalog no. R930-25)
<i>AOX1</i> Transcription Termination (TT) region	Native transcription termination and polyadenylation signal (~260 bp) from <i>AOX1</i> gene that permits efficient 3' mRNA processing, including polyadenylation, for increased mRNA stability
<i>TEF1</i> promoter (GenBank accession numbers D12478, D01130)	Transcription elongation factor 1 gene promoter from <i>Saccharomyces cerevisiae</i> that drives expression of the <i>Sh ble</i> gene in <i>Pichia</i> , conferring Zeocin [™] resistance
EM7 (synthetic prokaryotic promoter)	Constitutive promoter that drives expression of the <i>Sh ble</i> gene in <i>E. coli</i> , conferring Zeocin TM resistance
Sh ble gene (Streptoalloteichus hindustanus ble gene)	Zeocin [™] resistance gene (Calmels <i>et al.</i> , 1991; Drocourt <i>et al.</i> , 1990; Gatignol <i>et al.</i> , 1988).
<i>CYC1</i> transcription termination region (GenBank accession number M34014)	3' end of the <i>Saccharomyces cerevisiae CYC1</i> gene that allows efficient 3' mRNA processing of the <i>Sh ble</i> gene for increased stability
pUC origin	Allows replication and maintenance of the plasmid in <i>E. coli</i>



The pGAPZ and pGAPZ α vectors do not contain a yeast origin of replication. Transformants can only be isolated if recombination occurs between the plasmid and the *Pichia* genome.

pGAPZ A, B, and C

Map of pGAPZ A, B, and C

The figure below summarizes the features of pGAPZ A (2884 bp), pGAPZ B (2882 bp), and pGAPZ C (2883 bp) vectors. The complete nucleotide sequences for pGAPZ A, B, and C are available for downloading from our Web site (www.invitrogen.com) or from Technical Service (page 34). Details of the multiple cloning sites are shown on page 11 for pGAPZ A, page 12 for pGAPZ B, and page 13 for pGAPZ C.



pGAPZ α A, B, and C

B, and C

Map of pGAPZ α A, The figure below summarizes the features of pGAPZ α A (3147 bp), pGAPZ α B (3151 bp), and pGAPZ α C (3152 bp) vectors. The complete nucleotide sequences for pGAPZ α A, B, and C are available for downloading from our Web site (www.invitrogen.com) or from Technical Service (page 34). Details of the multiple cloning sites are shown on page 15 for pGAPZ α A, page 16 for pGAPZ α B, and page 17 for pGAPZ α C.



Methods

General Cloning Considerations

Introduction	Before cloning your gene into one of the pGAPZ or pGAPZ α vectors, please consider some of the general guidelines presented below. The multiple cloning sites for pGAPZ A, B, and C are located on pages 11-13 and the multiple cloning sites for pGAPZ α A, B, and C are presented on pages 15-17 to help you develop a cloning strategy.
General Considerations	The following are some general considerations applicable to pGAPZ A, B, and C and pGAPZ α A, B, and C.
	• The codon usage in <i>Pichia</i> is believed to be similar to <i>Saccharomyces cerevisiae</i> .
	• Many Saccharomyces genes have proven to be functional in Pichia.
	• Plasmid constructions should be maintained in a <i>recA</i> , <i>endA E</i> . <i>coli</i> strain such as TOP10F'.
	• You may wish to express your gene of interest without the C-terminal peptide. In this case, be sure your gene contains a stop codon.
	• The premature termination of transcripts because of "AT rich regions" has been observed in <i>Pichia</i> and other eukaryotic systems (Henikoff and Cohen, 1984; Irniger <i>et al.</i> , 1991; Scorer <i>et al.</i> , 1993; Zaret and Sherman, 1984). If you have problems expressing your gene, check for premature termination by northern analysis and check your sequence for AT rich regions. It may be necessary to change the sequence in order to express your gene (Scorer <i>et al.</i> , 1993).
Special Considerations for pGAPZ	 Be sure your gene has an initiation ATG as part of a Kozak consensus sequence (Kozak, 1987; Kozak, 1990). The Kozak consensus sequence is: ANN<u>ATG</u>G
	• For expression of a fusion protein containing the <i>myc</i> epitope and the polyhistidine tag, the open reading frame (ORF) of your gene must be cloned in frame with the C-terminal peptide.
Special Considerations for	• The initiation ATG in the α -factor signal sequence corresponds to the native initiation ATG of the <i>GAP</i> gene.
pGAPZα	• The open reading frame (ORF) of the mature gene of interest should be cloned in frame and downstream of the α -factor signal sequence and in frame with the C-terminal tag (if desired).
	• The predicted protease cleavage sites for the α -factor signal sequence are indicated in the figures on pages 15-17.

General Cloning Considerations, continued



You will need to prepare competent E. coli cells for transformation. Please see Current Protocols in Molecular Biology (Ausubel et al., 1994) or Molecular Biology: A Laboratory Manual (Sambrook et al., 1989) for preparation of electrocompetent or chemically competent E. coli or use your laboratory's procedure.

We recommend that you transform pGAPZ A, B, and C and pGAPZ a, B, and C into a recA, endA E. coli strain (e.g. TOP10F') so that you have a permanent stock and are able to propagate the plasmids. Electrocompetent TOP10F' cells are available from Invitrogen.

Item	Amount	Catalog no.
One Shot® TOP10F'	21 x 50 µl (1.0 ml total)	C3030-03
TOP10F′ Ultracomp [™]	5 x 300 µl (1.5 ml total)	C665-03
TOP10F' Electrocomp [™]	5 x 80 µl (400 µl total)	C665-55
TOP10F' Electrocomp [™]	10 x 80 µl (800 µl total)	C665-11

Propagation of It is recommended that pGAPZ and pGAPZ α be transformed into an *E. coli* cell line for maintainence of the vector. pGAPZ and pGAPZα The pGAPZ and pGAPZ α plasmids are supplied lyophilized.

- 1 Add 20 μ l sterile water or TE buffer to the lyophilized vector to prepare a 1 μ g/ μ l stock solution.
- 2. Dilute 1 μ l of the plasmid (1 μ g/ μ l) to 10-100 pg/ μ l using sterile water or TE buffer.
- Transform competent E. coli with 1-2 µl of the diluted plasmid and select on Low 3 Salt LB with 25 µg/ ml Zeocin[™] (see recipe on page 18).
- 4. Isolate a single colony containing the plasmid.
- 5. Culture in 5 ml Low Salt LB with 25 µg/ml Zeocin[™]. Grow overnight at 37°C.
- 6. Mix thoroughly 0.85 ml of culture with 0.15 ml sterile glycerol.
- 7. Transfer to a freezer vial and freeze in liquid nitrogen or a dry ice/ethanol bath.
- 8. Store at -80°C.



For optimal activity of Zeocin[™], the salt concentration of the medium must remain low (< 90 mM) and the pH must be 7.5. Prepare Low Salt LB broth and plates using the recipe on page 16.

Failure to lower the salt content of your LB medium will result in non-selection due to inactivation of the drug.

Cloning Detailed maps of the multiple cloning sites are provided on the following pages. Please refer to Ausubel, et al., 1994, pages 3.16.1 to 3.17.3. or Sambrook, et al., 1989, pages **Procedures** 5.10 to 5.13. for general help with cloning.

Cloning into pGAPZ

Multiple Cloning Site of pGAPZ A		elow is the multiple e cleavage site. Th <i>pa</i> I site is unique t quencing and func	e cloning site of p e boxed nucleotic o pGAPZ A. The tional testing.	GAPZ A. Restri des indicate the v multiple cloning	iction sites are la variable region. F g site has been co	beled to indicate Please note that the onfirmed by
371	AACCACCA	GA ATCGAATATA	AAAGGCGAAC	ACCTTTCCCA	ATTTTGGTTT	CTCCTGACCC
			pG	AP forward priming	site	Sfu I
431	AAAGACTT	TA AATTTAATTT	ATTTGTCCCT	ATTTCAATCA	ATTGAACAAC	TATTTCGAAA
491	EcoR I CGAGGAAT	<i>PmI</i> I IC ACGTGGCCCA	Sf/ I GCCGGCCGTC	Asp718 I TCGGATCGGT	Kpn Xho 	Sac II Not I
551	CCAGCTTG	Apal GG CCC GAA CA Glu Gl	A AAA CTC A' n Lys Leu II	<i>nyc</i> epitope IC TCA GAA (Le Ser Glu (GAG GAT CTG Glu Asp Leu	AAT AGC GCC Asn Ser Ala
603	GTC GAC Val Asp	Polyhistidine CAT CAT CAT C His His His H	tag CAT CAT CAT Lis His His	IGA GTTTTAG(* * *	CCT TAGACATO	GAC TGTTCCTCAC
660	TTCAAGTT	GG GCACTTACGA	GAAGACCGGT	CTTGCTAGAT	TCTAATCAAG	AGGATGTCAG
720	3´ AOX1 prin AATGCCAT	IT GCCTGAGAG	A TGCAGGCTT	C ATTTTTGAT	A CTTTTTTAT:	Г ТСТААССТАТ

780 ATAGTATAGG ATTTTTTTG TCATTTTGTT TCTTCTCGTAC

Cloning into pGAPZ, continued

Multiple C Site of pG	loning APZ B	Below is the multip the cleavage site. T <i>Xba</i> I site is unique sequencing and fun	ble cloning site of the boxed nucleo to pGAPZ B. The totional testing.	f pGAPZ B. Rest tides indicate the ne multiple clonin	riction sites are l variable region. ng site has been	abeled to indicate Please note that the confirmed by
371	AACCACCA	GA ATCGAATATA	AAAGGCGAAC	ACCTTTCCCA	ATTTTGGTTT	CTCCTGACCC
			pG	AP forward priming	site	Sfu I
431	AAAGACTT	TA AATTTAATTT	ATTTGTCCCT	ATTTCAATCA	ATTGAACAAC	TATTTCGAAA
	EcoR I	Pml I	Sfl I	Asp7181	Kpn IXho I	Sac II Not I
491	CGAGGAAT	IC ACGTGGCCCA	GCCGGCCGTC	TCGGATCGGT	ACCTCGAGCC	GCGGCGGGCCG
551	X] CCAGCTT	bal <u>TCTA</u> GAA CAA J Glu Gln	<i>my</i> AAA CTC ATC Lys Leu Ile	<mark>cepitope</mark> TCA GAA GAG Ser Glu Glu	G GAT CTG AA 1 Asp Leu As	AT AGC GCC Sn Ser Ala
601	GTC GAC	Polyhistidine t CAT CAT CAT C. His His His H	tag AT CAT CAT is His His	IGA GTTTTAGC ***	CCT TAGACATO	GAC TGTTCCTCAG
658	TTCAAGTT	GG GCACTTACGA	GAAGACCGGT	CTTGCTAGAT	TCTAATCAAG	AGGATGTCAG
	3´ AOX1 prim	ing site				
718	AATGCCA'	ITT GCCTGAGAG	A TGCAGGCTTO	C ATTTTTGATA	Α СТТТТТТТАТ]	TGTAACCTAT

778 ATAGTATAGG ATTTTTTTG TCATTTTGTT TCTTCTCGTA CGAGCTTG

Cloning into pGAPZ, continued

Multiple Clo Site of pGA	oning PZ C	Belov the cl <i>Sna</i> B seque	v is the multiple eavage site. The I site is unique ncing and function	cloning site of p boxed nucleotid to pGAPZ C. Th ional testing.	GAPZ C. Restrie les indicate the v e multiple clonir	ction sites are lal ariable region. P ng site has been c	beled to indicate lease note that the confirmed by
371	AACCACO	CAGA	ATCGAATATA	AAAGGCGAAC	ACCTTTCCCA	ATTTTGGTTT	CTCCTGACCC
				pG	AP forward priming	site	Sfu I
431	AAAGACI	TTTA	AATTTAATTT	ATTTGTCCCT	ATTTCAATCA	ATTGAACAAC	TATTTCGAAA
491	EcoR CGAGGAA	I ATTC	<i>Pml</i> I ACGTGGCCCA	<i>Sfl</i> I GCCGGCCGTC	Asp718 TCGGATCGGT	Kpn Xho - ACCTCGAGCC	Sac II Not I
551	CCAGCTI	SnaE ACG	GAA CAA Glu Gln	AAA CTC ATC Lys Leu Ile	nycepitope C TCA GAA GA e Ser Glu G	AG GAT CTG lu Asp Leu .	AAT AGC GCC Asn Ser Ala
602	GTC GAG Val As	C CAI p His	Polyhistidine t CAT CAT CA S His His H	tag AT CAT CAT ' is His His	IGA GTTTTAG(***	CCT TAGACAT	GAC TGTTCCTCAG
659	TTCAAG	TTGG	GCACTTACGA	GAAGACCGGT	CTTGCTAGAT	TCTAATCAAG	AGGATGTCAG
	3´ AOX1 p	riming	site				
719	AATGCCA	ATTT	GCCTGAGAGA	TGCAGGCTTC	ATTTTTGATA	CTTTTTTATT	TGTAACCTAT
779	ATAGTAI	TAGG	ATTTTTTTG	TCATTTTGTT	TCTTCTCGTA	CGAGCTT	

Cloning into $pGAPZ\alpha$

Introduction	The pGAPZ α vectors possess the α -factor mating signal sequence for secretion of your protein. The information in this section is provided to assist you in designing a cloning strategy. Details of the multiple cloning sites of the pGAPZ α A, B, and C vectors can be found on the following pages.
Cloning Considerations	The variable region of pGAPZ α A, B, and C is located between the α -factor signal sequence and the multiple cloning site. It is designed to facilitate cloning into the vector in frame with the signal sequence . Note that if you wish to create a fusion protein with the C-terminal tag, you must consider the 3' cloning sequence as well. If your insert does not conveniently ligate in frame with both terminal fusion partners, it may be necessary to use PCR to create ends that will result in in-frame fusions. Please consider both the frame of the signal sequence and the C-terminal fusion when designing a cloning strategy.
Signal Sequence Processing	 The processing of the α-factor mating signal sequence in pGAPZα occurs in two steps: The preliminary cleavage of the signal sequence by the <i>KEX2</i> gene product, with the final Kex2 cleavage occurring between arginine and glutamine in the sequence Glu-Lys-Arg * Glu-Ala-Glu-Ala, where * is the site of cleavage. The Glu-Ala repeats are further cleaved by the <i>STE13</i> gene product.
Optimization of Signal Cleavage	In <i>Saccharomyces cerevisiae</i> , it has been noted that the Glu-Ala repeats are not necessary for cleavage by Kex2, but cleavage after Glu-Lys-Arg may be more efficient when followed by Glu-Ala repeats. A number of amino acids are tolerated at site X instead of Glu in the sequence Glu-Lys-Arg-X. These amino acids include the aromatic amino acids, small amino acids, and histidine. Proline, however, will inhibit Kex2 cleavage. For more information on Kex2 cleavage, please see (Brake <i>et al.</i> , 1984)
	There are some cases where Ste13 cleavage of Glu-Ala repeats is not efficient, and Glu- Ala repeats are left on the N-terminus of the expressed protein of interest. This is generally dependent on the protein of interest.
Cloning Your Gene Flush with the Kex2 Cleavage Site	If you wish to have your protein expressed with a native N-terminus, you may use the <i>Xho</i> I site at bp 736-741 to clone your gene flush with the Kex2 cleavage site. Use PCR to rebuild the sequence from the <i>Xho</i> I site to the arginine codon at nucleotides 745-747. Remember to include the first amino acid(s) of your protein, if necessary, for correct fusion to the Kex2 cleavage site.

Cloning into $pGAPZ\alpha$, continued

Multiple Clo Site of pGA	ning PZα A	Below is the cleav cloning	the mul vage site. site has b	tiple c The b een co	loning oxed onfirm	g site o nucleo red by	of pG. otide seque	APZα indica encing	A. Re tes the g and f	estrict e varia functio	ion si able re onal te	tes are egion. esting	e labe The i	led to nultip	indicate le
361	GATTATT	gga aa	CCACCA	GA A1	rcga <i>p</i>	ATATA	A AAA	AGGCO	GAAC pGA	ACCT	TTTCC vard pri	CCA A	ATTTT ite	rggti	ГТ
421	CTCCTGA	CCC AA	AGACTT'	TA AA	ATTTA	ATTI	AT1	TGTC	CCCT	ATTI	CAAI	CA A	ATTGZ	AACAA	AC
481	TATTTCG	AAA CG	ATG A Met A	GA TI rg Pł	IT CC ne Pr	CT TO	CA Al er Il	TT TT .e Pł	TT AC ne Th	CT GO nr Al	CT GI La Va	TT TT al Le	FA TI eu Pi	TC GC ne Al	CA La
532	GCA TCC Ala Ser	TCC G Ser A	CA TTA la Leu	GCT Ala	GCT Ala	CCA Pro	GTC Val	AAC Asn	ACT Thr	ACA Thr	ACA Thr	GAA Glu	GAT Asp	GAA Glu	ACG Thr
					α-fa	actor si	gnal se	equenc	е						
583	GCA CAA Ala Gln	ATT C Ile P	CG GCT ro Ala	GAA Glu	GCT Ala	GTC Val	ATC Ile	GGT Gly	TAC Tyr	TCA Ser	GAT Asp	TTA Leu	GAA Glu	GGG Gly	GAT Asp
634	TTC GAT Phe Asp	GTT G Val A	CT GTT la Val	TTG Leu	CCA Pro	TTT Phe	TCC Ser	AAC Asn	AGC Ser	ACA Thr	AAT Asn	AAC Asn	GGG Gly	TTA Leu	TTG Leu
685	TTT ATA Phe Ile	AAT A Asn T	CT ACT hr Thr	ATT Ile	GCC Ala	AGC Ser	ATT Ile	GCT Ala	GCT Ala	AAA Lys	GAA Glu	GAA Glu	GGG Gly	GTA Val	TCT Ser
736	Xhol* CTC GAG Leu Glu	AAA A Lys A	GA GAG	GCT Ala	GAA Glu	GCT Ala	EcoR GAAI	I TTCAC	Pml I GTG	GCCC	SA GC	Sfil CCGGG	CCGT	С ТСС	GGATC
As	50718 Kon	Xho I	Sac II N	ot I	i s sign	iai ciea	xaye Xba	1			тус	epitop	e		
793	GGTACCT	CGA GC	CGCGGC	GG C(CGCCA	AGCTI	TCT po	TA GA GI	AA CA Lu GI line tag	AA AA ln Ly	AA CI ys Le	C Ar eu I	IC TO le Se	CA GA er Gi	AA GAG lu Glu
851	GAT CTG Asp Leu	AAT A Asn S	GC GCC er Ala	GTC Val	GAC Asp	CAT His	CAT His	CAT His	CAT His	CAT His	CAT His	TGA ***	GTT	ГТАGO	C CTTA
905	GACATGA	CTG TT	CCTCAG	TT CA	AAGTI	GGGG	C ACI	TACO	GAGA	AGAC	CCGGI	CT 1	rgct <i>i</i>	AGATI	ГС ТААТ
		3´ A	OX1 primi	ng site											
969	CAAGAGG	ATG TC	AGAATG	CC AI	TTTGC	CTG	A GAC	GATGO	CAGG	CTTC	CATTI	TT (GATA	СТТТЭ	FT TATT
1033	TGTAACC	TAT AT	AGTATA	GG AI	TTTT	TTTG	TCA	TTTT	GTT	ТСТІ	CTCG	Ţ			
*	The Xho I si cleavage site	te upstrear e (see pag	n of the Ke e 14).	ex2 clea	avage s	ite is u	sed to	clone t	he gen	e of int	erest fl	ush wi	th the k	Kex2	

Cloning into $pGAPZ\alpha$, continued

Multiple Cl Site of pG/	oning PZα B Below is the multiple cloning site of pGAPZ α B. Restriction sites are labeled to indicate the cleavage site. The boxed nucleotides indicate the variable region. Please not that the <i>Pst</i> I site is unique to pGAPZ α B. The multiple cloning site has been confirmed by sequencing and functional testing.
361	GATTATTGGA AACCACCAGA ATCGAATATA AAAGGCGAAC ACCTTTCCCA ATTTTGGTTT pGAP forward priming site
421	CTCCTGACCC AAAGACTTTA AATTTAATTT ATTTGTCCCT ATTTCAATCA ATTGAACAAC
481	TATTTCGAAA CG ATG AGA TTT CCT TCA ATT TTT ACT GCT GTT TTA TTC GCA Met Arg Phe Pro Ser Ile Phe Thr Ala Val Leu Phe Ala
532	GCA TCC TCC GCA TTA GCT GCT CCA GTC AAC ACT ACA ACA GAA GAT GAA ACG Ala Ser Ser Ala Leu Ala Ala Pro Val Asn Thr Thr Glu Asp Glu Thr
	α-factor signal sequence
583	GCA CAA ATT CCG GCT GAA GCT GTC ATC GGT TAC TCA GAT TTA GAA GGG GAT Ala Gln Ile Pro Ala Glu Ala Val Ile Gly Tyr Ser Asp Leu Glu Gly Asp
634	TTC GAT GTT GCT GTT TTG CCA TTT TCC AAC AGC ACA AAT AAC GGG TTA TTG Phe Asp Val Ala Val Leu Pro Phe Ser Asn Ser Thr Asn Asn Gly Leu Leu
685	TTT ATA AAT ACT ACT ATT GCC AGC ATT GCT GCT AAA GAA GAA GGG GTA TCT Phe Ile Asn Thr Thr Ile Ala Ser Ile Ala Ala Lys Glu Glu Gly Val Ser
	Xho I* Rex2 signal cleavage Pst I EcoR I Pm/ I Sfi I
736	CTC GAG AAA AGA GAG GCT GAA GCT GCAGG AATTCACG TGGCCCAGC CGGCCGTCTC Leu Glu Lys Arg Glu Ala
	Asp718 Kpn Xho Sac Not Xha myc epitope
792	GGATCGGTAC CTCGAGCCGC GGCGGCCGCC AGCTTTCTA GAA CAA AAA CTC ATC TCA GAA Glu Gln Lys Leu Ile Ser Glu polyhistidine tag
852	GAG GAT CTG AAT AGC GCC GTC GAC CAT CAT CAT CAT CAT CAT TGA GTTTTAGCC Glu Asp Leu Asn Ser Ala Val Asp His His His His His ***
907	TAGACATGAC TGTTCCTCAG TTCAAGTTGG GCACTTACGA GAAGACCGGT CTTGCTAGAT 3´ AOX1 priming site
967	TCTAATCAAG AGGATGTCAG AATGCCATTT GCCTGAGAGA TGCAGGCTTC ATTTTTGATA
1027	CTTTTTTATT TGTAACCTAT ATAGTATAGG ATTTTTTTTG TCATTTTGTT TCTT * The <i>Xho</i> I site upstream of the Kex2 cleavage site is used to clone the gene of interest flush with the Kex2 cleavage site (see page 14).

Cloning into $pGAPZ\alpha$, continued

Multiple Cl Site of pGA	Derived Below is the multiple cloning site of pGAPZ α C. Restriction sites are labeled to indicate the cleavage site. The boxed nucleotides indicate the variable region. Please not that the <i>Cla</i> I site is unique to pGAPZ α C. The multiple cloning site has been confirmed by sequencing and functional testing.
361	GATTATTGGA AACCACCAGA ATCGAATATA AAAGGCGAAC ACCTTTCCCA ATTTTGGTTT
	pGAP forward priming site
421	CTCCTGACCC AAAGACTTTA AATTTAATTT ATTTGTCCCT ATTTCAATCA ATTGAACAAC
481	TATTTCGAAA CG ATG AGA TTT CCT TCA ATT TTT ACT GCT GTT TTA TTC GCA Met Arg Phe Pro Ser Ile Phe Thr Ala Val Leu Phe Ala
532	GCA TCC TCC GCA TTA GCT GCT CCA GTC AAC ACT ACA ACA GAA GAT GAA ACG
	Ala Ser Ser Ala Leu Ala Ala Pro Val Asn Thr Thr Glu Asp Glu Thr
	α-factor signal sequence
583	GCA CAA ATT CCG GCT GAA GCT GTC ATC GGT TAC TCA GAT TTA GAA GGG GAT
	Ala Gln Ile Pro Ala Glu Ala Val Ile Gly Tyr Ser Asp Leu Glu Gly Asp
634	TTC GAT GTT GCT GTT TTG CCA TTT TCC AAC AGC ACA AAT AAC GGG TTA TTG
685	TTT ATA AAT ACT ACT ATT GCC AGC ATT GCT GCT AAA GAA GAA GGG GTA TCT Phe Ile Asn Thr Thr Ile Ala Ser Ile Ala Ala Lys Glu Glu Gly Val Ser
	Xho I* Kex2 signal cleavage Cla I EcoR I Pml I Sfi I
736	CTC GAG AAG AGA GAG GCT GAA GCA TCGAT GAATTCACGT GGCCCAGCCG GCCGTCT Leu Glu Lys Arg Glu Ala Glu Ala Ste13 signal cleavage
	Asp718 I Kpn Xho I Sac II Not I Xba I myc epitope
792	CGGATCGGTA CCTCGAGCCG CGGCGGCCGC CAGCTTTCTA GAA CAA AAA CTC ATC TCA
	polyhistidine tag
850	GAA GAG GAT CTG AAT AGC GCC GTC GAC CAT CAT CAT CAT CAT TGA GTT
	Glu Glu Asp Leu Asn Ser Ala Val Asp His His His His His ***
902	TAGCCTTAGA CATGACTGTT CCTCAGTTCA AGTTGGGCAC TTACGAGAAG ACCGGTCTTG
	3' AOX1 priming site
962	CTAGATTCTA ATCAAGA ['] GGA TGTCAGAATG CCATTTGC ['] CT GAGAGATGCA GGCTTCATTT
1022	TTGATACTTT TTTATTTGTA ACCTATATAG TATAGGATTT TTTTTGTCAT TTTGTTTCTT
	* The <i>Xho</i> I site upstream of the Kex2 cleavage site is used to clone the gene of interest flush with the Kex2 cleavage site (see page 14).

Transformation into E. coli

Introduction	Ligation mixtures may be transformed into <i>E. coli</i> and selected on Low Salt LB medium (see below) with Zeocin TM . Transformants are isolated and analyzed for the presence and orientation of insert. There is no blue/white screening for the presence of insert with pGAPZ or pGAPZ α . After obtaining the desired recombinant plasmid, transform the linearized construct into <i>Pichia</i> as described on page 20.						
Note	For Zeocin TM to be active, the salt concentration of the medium must remain low $(< 90 \text{ mM})$ and the pH must be 7.5. Prepare Low Salt LB broth and plates using the following recipe.						
	Failure to lower the salt content of your LB medium will result in non-selection due to inactivation of the drug.						
	Low Salt LB Medium:						
	10 g Tryptone 5 g NaCl 5 g Yeast Extract						
	 Combine the dry reagents above and add deionized, distilled water to 950 ml. Adjust pH to 7.5 with 1 N NaOH. Bring the volume up to 1 liter. For plates, add 15 g/L agar before autoclaving. 						
	2. Autoclave on liquid cycle at 15 lbs/sq. in. and 121°C for 20 minutes.						
	3. Allow the medium to cool to at least 55°C before adding the Zeocin [™] to 25 μg/ml final concentration.						
	4. Store plates at 4°C in the dark. Plates containing Zeocin [™] are stable for 1-2 weeks.						
E. coli	Guidelines are as follows:						
Transformation	• Transformation may be done by either electroporation or chemical methods. Use your preferred method or refer to general molecular biology references (Ausubel <i>et al.</i> , 1994; Sambrook <i>et al.</i> , 1989).						
	• Add either Low Salt LB or LB medium to the cells after heat shock or electroporation to allow them to recover.						
	 Plate on Low Salt LB medium with 25 µg/ml Zeocin[™]. Note: You may also use SOB, 2XYT, or TB medium, but you may have to increase the concentration of Zeocin[™] to 50 µg/ml to compensate for differences in the salt concentration. 						
	• Incubate overnight at 37°C.						

Transformation into *E. coli*, continued

Analysis of Transformants	 After transformation, LB plates with 25 μg/ resistant colonies. Pick 10 Zeocin[™]-resis Salt LB medium with Isolate plasmid DNA below). Be sure to make a gly Purify the clone by Prepare an overnigl Combine 0.85 ml o Mix by vortexing at Freeze the tube in 11 Store at -80°C 	plate 10 µl and 100 µl of the transformation mix onto Low Salt ml Zeocin [™] (see Recipe , previous page) and select Zeocin [™] - stant transformants and inoculate each colony into 2 ml Low 25 µg/ml Zeocin [™] . Grow overnight at 37°C with shaking. by miniprep for restriction analysis and sequencing (see cerol stock of your purified clone for safekeeping. streaking for single colonies at bacterial culture from a single colony f the overnight bacterial culture with 0.15 ml of sterile glycerol nd transfer to a labeled storage tube iquid nitrogen or a dry ice/ethanol bath
Recombinant Clones	<i>Pichia</i> to confirm that your C-terminal tag. To sequenc using the pGAP Forward an primers are not available se placing custom primer orde Custom Primers.	gene is in frame with the α -factor secretion signal and/or the e your construct in pGAPZ or pGAPZ α , we recommend ad the 3' <i>AOX1</i> primer sequences. Please note that these eparately from Invitrogen. If you would like information on rrs, visit our Web site (www.invitrogen.com) and select
	Sequencing Primer	Sequence
	pGAP Forward	5'-GTCCCTATTTCAATCAATTGAA-3'
	3' AOX1	5'-GCAAATGGCATTCTGACATCC-3'
	For sequencing protocols, p Biology (Ausubel et al., 19 Manual (Sambrook et al., 1	blease refer to Unit 7 in <i>Current Protocols in Molecular</i> 94) or Chapter 13 in <i>Molecular Cloning: A Laboratory</i> 989).
Plasmid Preparation	Once you have cloned and transform <i>Pichia</i> (5-10 µg o S.N.A.P. [™] Miniprep Kit (K you have purified plasmid l	sequenced your insert, generate enough plasmid DNA to of each plasmid per each transformation). We recommend the (1900-01) for quick purification of pure plasmid DNA. Once DNA, proceed to <i>Pichia</i> Transformation, next page.

Pichia Transformation

Introduction	At this point, you should have your gene cloned into one of the pGAPZ or pGAPZ α vectors. Your construct should be correctly fused to the α -factor secretion signal and/or C-terminal peptides. You will then prepare 5-10 µg of plasmid DNA and linearize the plasmid prior to transformation and selection in <i>Pichia</i> . Transformants are plated on YPDS plates containing 100 µg/ml Zeocin TM to isolate Zeocin TM -resistant clones.					
Method of Transformation	We do not recommend spheroplasting for transformation of <i>Pichia</i> with plasmids containing the Zeocin [™] resistance marker. Spheroplasting involves removal of the cell wall to allow DNA to enter the cell. Cells must first regenerate the cell wall before they are able to express the Zeocin [™] resistance gene. Plating spheroplasts directly onto selective medium containing Zeocin [™] will result in complete cell death.					
	We recommend electroporation for transformation of <i>Pichia</i> with the pGAPZ or pGAPZ α vectors. Electroporation yields 10 ³ to 10 ⁴ transformants per µg of linearized DNA and does not destroy the cell wall of <i>Pichia</i> . If you do not have access to an electroporation device, use the LiCl protocol on page 33 or the EasyComp TM Kit (Catalog no. K1730-01).					
Before Starting	You will need the following reagents for transforming <i>Pichia</i> and selecting transformants on $\text{Zeocin}^{\text{TM}}$. Note : Inclusion of sorbitol in YPD plates stabilizes electroporated cells as they appear to be somewhat osmotically sensitive.					
	• 5-10 μ g pure pGAPZ or pGAPZ α containing your insert					
	YPD Medium 50 ml conical polymonylong types					
	 I liter cold (+4°C) sterile water (place on ice the day of the experiment) 					
	 25 ml cold (+4°C) sterile 1 M sorbitol (place on ice the day of the experiment) 30°C incubator 					
	Electroporation device and 0.2 cm cuvettes					
	• YPDS plates containing 100 µg/ml Zeocin [™] (please see page 30 for recipe)					
Restriction Digest	1. Digest ~5-10 μg of plasmid DNA with either <i>Avr</i> II (191 bp) or <i>Bsp</i> HI (356 bp). Each enzyme cuts once in the <i>GAP</i> promoter region to linearize the vector. Choose the enzyme that does not cut within your gene.					
	2. We recommend that you check a small aliquot of your digest by agarose gel electrophoresis for complete linearization.					
	3. If the vector is completely linearized, heat inactivate or add EDTA to stop the reaction, phenol/chloroform extract once, and ethanol precipitate using 1/10 volume 3 M sodium acetate and 2.5 volumes of 100% ethanol.					
	4. Centrifuge the solution to pellet the DNA, wash the pellet with 80% ethanol, air-dry, and resuspend in 10 μ l sterile, deionized water. Use immediately or store at -20°C.					

Pichia Transformation, continued

Preparation of <i>Pichia</i> for	1.	Grow your <i>Pichia pastoris</i> strain in 5 ml of YPD in a 50 ml conical at 30°C overnight.
Electroporation	2.	Inoculate 500 ml of fresh medium in a 2 liter flask with 0.1-0.5 ml of the overnight culture. Grow overnight again to an $OD_{600} = 1.3-1.5$.
	3.	Centrifuge the cells at 1500 x g for 5 minutes at $+4^{\circ}$ C. Resuspend the pellet with 500 ml of ice-cold (0°C), sterile water.
	4.	Centrifuge the cells, then resuspend the pellet with 250 ml of ice-cold (0°C), sterile water.
	5.	Centrifuge the cells, then resuspend the pellet in 20 ml of ice-cold (0°C) 1 M sorbitol.
	6.	Centrifuge the cells, then resuspend the pellet in 1 ml of ice-cold (0°C) 1 M sorbitol for a final volume of approximately 1.5 ml. Keep the cells on ice and use that day. Do not store cells.
Transformation by Electroporation	1.	Mix 80 μ l of the cells from Step 6 (above) with 5-10 μ g of linearized DNA (in 5-10 μ l sterile water) and transfer them to an ice-cold (0°C) 0.2 cm electroporation cuvette.
	2.	Incubate the cuvette with the cells on ice for 5 minutes.
	3.	Pulse the cells according to the parameters for yeast (<i>Saccharomyces cerevisiae</i>) suggested by the manufacturer of the specific electroporation device being used.
	4.	Immediately add 1 ml of ice-cold 1 M sorbitol to the cuvette. Transfer the cuvette contents to a sterile 15 ml tube.
	5.	Let the tube incubate at 30°C without shaking for 1 to 2 hours.
	6.	Spread 10, 25, 50, 100, and 200 μ l each on separate, labeled YPDS plates containing 100 μ g/ml Zeocin TM . Plating at low cell densities favors efficient Zeocin TM selection.
	7.	Incubate plates for 2 to 3 days at 30°C until colonies form.
	8.	Pick 10-20 colonies and purify (streak for single colonies) on fresh YPD or YPDS plates containing 100 µg/ml Zeocin [™] .

Pichia Transformation, continued



Generally several hundred to several thousand ZeocinTM-resistant colonies are generated using the above protocol. If more colonies are needed the protocol may be modified as described below. Please note that you will need ~20 150 mm plates with YPDS agar containing 100 μ g/ml ZeocinTM.

- 1. Set up two transformations per construct and follow Steps 1 through 5 of the **Transformation by Electroporation** protocol, previous page.
- 2. After 1 hour in 1 M sorbitol at 30°C (Step 5, previous page), add 1 ml YPD medium to each tube.
- 3. Shake (~ 200 rpm) the cultures at 30°C.
- 4. After 1 hour, take one of the tubes and plate out all of the cells by spreading 200 μl on 150 mm plates containing 100 μg/ml Zeocin[™].
- (Optional) Continue incubating the other culture for three more hours (for a total of four hours) and then plate out all of the cells by spreading 200 µl on 150 mm plates containing 100 µg/ml Zeocin[™].
- 6. Incubate plates for 2 to 4 days at 30°C until colonies form.

Analysis of *Pichia* Transformants

If you wish, you may analyze 6-10 of your ZeocinTM-resistant *Pichia* transformants for the presence of insert using PCR, or for copy number using Southern analysis. Please refer to a *Pichia* Expression Kit manual for these protocols.

If you choose to analyze for the presence of insert using PCR, please note that the parent plasmids will produce the following sized PCR products when using the pGAP Forward and the 3' *AOX1* primer sequences:

Vector	PCR Product
pGAPZ	~275 bp
pGAPZα	~540 bp

Remember to add the size of these fragments to the size of your insert to interpret your PCR results.

You are now ready to test your transformants for expression of your gene. Please see **Expression in** *Pichia*, next page.

Isolation of Multicopy Recombinants *in vivo*

A quick, direct way to select putative multi-copy recombinants is to plate the transformation mix on increasing concentrations of $\text{Zeocin}^{\text{TM}}$. Please note that multiple gene insertion with the pGAPZ and pGAPZ α has not been functionally tested by Invitrogen.

- Prepare YPDS plates containing 500, 1000, and 2000 µg/ml Zeocin[™]
- Plate 100 to 200 μl of the transformation mix on each plate and incubate at 30°C for 2 days



It is strongly recommended that you purify Zeocin[™] resistant transformants at least once. The following steps for expression do not utilize Zeocin[™] selection and any mixed colony transformants may lose the expressing clone.

- Streak out the chosen Zeocin[™]-resistant *Pichia* transformant on YPD plates containing Zeocin[™].
- Grow for 2-4 days at 30°C until single colonies form

Expression in Pichia

Introduction	The primary purpose of small-scale expression is to identify/confirm a recombinant <i>Pichia</i> clone that is expressing the correct protein. Small-scale expression conditions may not be optimal for your protein. For this reason, the method you choose for detection (i.e. SDS-PAGE, western, or functional assay) may be an important factor in determining the success of expression. If your method of detection does not reveal any expression, you may want to consider using a more sensitive detection method. Once a positive clone has been identified, large scale expression can be carried out in shake flask or fermentation and expression conditions optimized.
Expression Guidelines	Expression can be done in either YPD medium (1% yeast extract, 2% peptone, 2% glucose) or Yeast Nitrogen Base with 0.5% glucose.
	1. Using a single colony, inoculate 10 ml of YPD. Grow at 28-30°C in a shaking incubator (250-300 rpm) overnight.
	2. Use 0.1 ml of the overnight culture to inoculate 50 ml of YPD in a 250 ml flask. Grow at 28-30°C in a shaking incubator (250-300 rpm).
	3. At each of the times indicated below transfer 1 ml of the expression culture to a 1.5 ml microcentrifuge tube. These samples will be used to analyze expression levels and determine the optimal time to harvest. Centrifuge at maximum speed in a tabletop microcentrifuge for 2-3 minutes at room temperature.
	Time points (hours): 0, 24 (1 day), 48 (2 days), 72 (3 days), 96 (4 days)
	4. For secreted expression, transfer the supernatant to a separate tube. Store the supernatant and the cell pellets at -80°C until ready to assay. Freeze quickly in liquid N_2 or a dry ice/ethanol bath.
	For intracellular expression, decant the supernatant and store just the cell pellets at -80°C until ready to assay. Freeze quickly in liquid N_2 or a dry ice/ethanol bath.

Expression in Pichia, continued

Detection of Recombinant Proteins in *Pichia*

We recommend that you use the following techniques to analyze the expression of your protein. Analyze BOTH the cells and the medium for the presence of your recombinant protein. Please note that the α -factor signal sequence (in pGAPZ α) will add approximately 9.3 kDa to your protein if it is unprocessed. The C-terminal tag will add 2.5 kDa to your protein. Be sure to account for any additional amino acids that are in between the signal sequence processing sites and the N-terminus of your protein (in pGAPZ α) and also the end of your protein and the C-terminal tag.

Technique	Method of Detection	Sensitivity
SDS-PAGE	Visualization by eye	Can detect as little as 100 ng
(Coomassie-stained)		in a single band
SDS-PAGE	Visualization by eye	Can detect as little as 2 ng in a
(Silver-stained)		single band
Western Analysis	Antibody to your particular protein Anti- <i>myc</i> Antibody (Catalog no. R950-25) Anti-His(C-term) Antibody (catalog no. R930-25)	Can detect as little as 1-10 pg, depending on detection method (alkaline phosphatase, horseradish peroxidase, radiolabeled antibody)
Functional assay	Varies depending on assay	Varies depending on assay
		Used to compare relative amounts of protein.

Scale-up of Expression

You may want to scale-up your expression protocol to produce more protein. This may be done by increasing the culture volume using larger baffled flasks or fermentation (call Invitrogen Technical Service, see below).



Because the pGAPZ and pGAPZα vectors do not contain the *HIS4* gene, *his4 Pichia* strains containing the integrated plasmid must be grown in complex medium (i.e. YPD) or minimal medium containing 0.004% histidine (available from Invitrogen, Catalog no. Q300-23). If histidine is not present in the medium, the cells will not grow.

Optimization of Pichia Protein Expression

Introduction	If you obtain no or low protein expression in your initial expression experiment, it may be necessary to optimize expression. Please refer to the following guidelines for assistance. Remember that <i>GAP</i> promoter is a constitutively expressed promoter. If the gene product is toxic to the cells, you may have to try an inducible expression system.
Proteolysis or Degradation	 Do a time course study of expression. Check to see if there is a time point that yields a larger percentage of full-length protein. If the protein is being degraded, try expressing in a protease deficient strain like SMD1168 (Catalog no. C175-00).
Low Secreted Expression Levels	 Check the cell pellet to see if overall expression is low or if the protein did not secrete. If it did not secrete, try a different signal sequence (e.g. a native signal sequence). Concentrate your supernatant by ammonium sulfate precipitation or ultrafiltration. Try expression with a higher density culture.
Low Expression Levels	 Look for multi-copy recombinants (i.e. jackpot clones) by slot blot (please refer to a <i>Pichia</i> Expression manual). There are many examples of increasing the expression levels of a particular protein by increasing the gene dosage (Clare <i>et al.</i>, 1991a; Clare <i>et al.</i>, 1991b; Romanos <i>et al.</i>, 1991). Scale up to fermentation. <i>Pichia</i> is particularly well suited to fermentation. Please call Invitrogen Technical Service for recommendations (page 34).
No Expression	Be sure to try some of the easier things listed above as no expression can be the same thing as very low expression. If none of these things improve protein expression, perform a northern blot analysis to check for transcription of your gene. Refer to a <i>Pichia</i> Expression Kit manual for a protocol for RNA isolation. If you see premature transcriptional termination, check the AT content of your gene. In <i>Saccharomyces</i> , there are a few consensus sequences which promote premature termination. One of these, TTTTTATA, resembles a sequence in the HIV-1 gp120 gene, ATTATTTTATAAA, which when expressed in <i>Pichia</i> showed premature termination of the mRNA. When this sequence was changed, longer transcripts were found (Scorer <i>et al.</i> , 1993).
Hyperglycosyla- tion	 If your protein is hyperglycosylated: Try intracellular expression. Your protein will not go through the secretion pathway and therefore, will not be modified. Deglycosylate the protein by treatment with Peptide:N-Glycosidase F or refer to a <i>Pichia</i> Expression Kit manual for other enzymes. Engineer the gene to remove any N-glycosylation sites (Asn-X-Ser/Thr).

Purification

Introduction	In this section, you will grow and induce a 10-200 ml culture of your <i>Pichia</i> transformant for trial purification on a metal-chelating resin such as ProBond TM . You may harvest the cells and store both the supernatant (medium) and the cells at -80°C until you are ready to purify your fusion protein, or you may proceed directly with protein purification. Please note that this section only describes sample application onto ProBondTM . For instructions on how to prepare and use ProBond TM resin, please refer to the ProBond TM Purification manual.	
ProBond [™] Resin	We recommend that you use the ProBond [™] Purification System (Catalog no. K850-01) for purifying fusion proteins expressed from pGAPZ or pGAPZα. Please note that instructions for equilibration of and chromatography on ProBond [™] resin are contained in the ProBond [™] Purification System Kit.	
	If you are using a metal-chelating resin other than ProBond [™] , please follow the manufacturer's recommendations for fusion proteins expressed in bacteria or yeast.	
Binding Capacity of ProBond [™]	One milliliter of ProBond [™] resin binds at least 1 mg of recombinant protein. This amount can vary depending on the protein.	
Q Important	Throughout the following protocol, be sure to keep the medium and fractions on ice. Small-scale purifications using the 2 ml ProBond [™] columns and buffers can be done at room temperature on the bench top. For large scale purifications, all reagents must be at +4°C.	
Expression of Secreted Protein	Express your protein using a small-scale culture and the optimal conditions for expression (if determined). Once your protein is expressed, separate the cells from the medium by centrifugation. Store the medium at -80°C or proceed directly to purification. If desired, the cells can be stored at -80°C for future analysis.	

Purification, continued

Preparation of Cell Lysates	Express your protein using a small-scale culture and the optimal conditions for expression (if determined). Once your protein is expressed, follow the protocol below to prepare a cell lysate for chromatography on ProBond [™] .		
	Prepare Breaking Buffer (BB) as described in the Recipes, page 30.		
	1.	Wash cells once in BB by resuspending them and centrifuging 5-10 minutes at $3000 \text{ x g at } +4^{\circ}\text{C}$.	
	2.	Resuspend the cells to an OD_{600} of 50-100 in BB.	
	3.	Add an equal volume of acid-washed glass beads (0.5 mm). Estimate volume by displacement.	
	4.	Vortex the mixture for 30 seconds, then incubate on ice for 30 seconds. Repeat 7 more times. Alternating vortexing with cooling keeps the cell extracts cold and reduces denaturation of your protein.	
	5.	Centrifuge the sample at +4°C for 5-10 minutes at 12,000 x g.	
	6.	Transfer the clear supernatant to a fresh container and analyze for your protein. The total protein concentration should be around 2-3 mg/ml.	
	7.	Save the pellet and extract with 6 M urea or 1% Triton X-100 to check for insoluble protein.	
Sample Applica- tion (Native Conditions)	The following protocol can be used for chromatography of medium. For sample application onto ProBond [™] , you will need Native Binding Buffer, pH 7.8 and a 2 ml ProBond [™] column, pre-equilibrated using native conditions.		
,	1.	Combine 1 ml of medium with 7 ml Native Binding Buffer.	
	2.	Take a pre-equilibrated ProBond [™] column and resuspend the resin in 4 ml of the diluted medium from Step 1.	
	3.	Seal the column and batch-bind by rocking gently at room temperature for 10 minutes.	
	4.	Let the resin settle by gravity or low speed centrifugation (800 x g) and carefully remove the supernatant. Save the supernatant to check for unbound protein.	
	5.	Repeat Steps 2 through 4 with the remaining 4 ml of diluted medium. Proceed to Column Washing and Elution Under Native Conditions in the ProBond [™] Purification manual. Use the recommendations noted for bacterial cell lysates.	
Sample Applica- tion (Denaturing Conditions)	Use Bin Buf	the protocol above except pre-equilibrate the ProBond [™] column using Denaturing ding Buffer and combine 1 ml of the medium with 7 ml of the Denaturing Binding fer.	
Note	We usin Puri Pici	have observed that some <i>Pichia</i> proteins may be retained on the ProBond [™] column ng native purification conditions. Optimization of the purification (see ProBond [™] ification manual) or using denaturing purification may remove these non-specific <i>hia</i> proteins.	

Purification, continued

Analysis of Purification	Be sure to save all fractions, washes, and flow-through for analysis by SDS-PAGE. You may need to use western blot analysis to detect your protein if expression is low or not enough protein was loaded onto the column. Please refer to the ProBond [™] Purification System manual for a guide to troubleshoot chromatography.
Scale-up	You may find it necessary to scale-up your purification to obtain sufficient amounts of purified protein. Adjust the pH and NaCl concentration of the medium with 1/10 volume of 10X Stock Solution B (ProBond TM Purification System) before adding it to the column. The pH should be \geq 7.5 and the NaCl concentration should be \sim 500 mM. Using 10X Stock Solution B to adjust the pH and the ionic strength keeps the total volume small for sample application.

Appendix

Recipes				
Yeast Nitrogen Base	Yeast Nitrogen Base is available separately from Invitrogen. Please see the table below for ordering information.			
	Item	Amount	Catalog no.	
	Yeast Nitrogen Base (YNB)	1 x 67 g pouch	Q300-07	
	-with ammonium sulfate	(each pouch contains reagents to		
	-without amino acids	prepare 500 ml of a 10X YNB solution)		
		500 g	Q300-09	
YPD or YEPD	Yeast Extract Peptone Dextrose Medium (1 liter) 1% yeast extract 2% peptone 2% dextrose (D-glucose)			
	Note: If you are using the YP Base Medium or the YP Base Agar medium pouches, follow the directions on the pouch.			
	1. Dissolve 10 g yeast extract and 20 g of peptone in 900 ml of water. Note : Add 20 g of agar if making YPD slants or plates.			
	2. Autoclave for 20 minutes on liquid cycle.			
	3. Add 100 ml of 20% Dextrose.			
	The liquid medium is stored at room temperature. YPD slants or plates are stored at +4°C. The shelf life is several months.			
YPD (+ Zeocin™)	Yeast Extract Peptone Dextrose Medium (1 liter)			
	1% yeast extract 2% peptone 2% dextrose (D-glucose) ± 2% agar ± 100 μg/ml Zeocin [™]			
	1. Dissolve the following in 900 ml of water:			
	10 g yeast extract 20 g of peptone			
	2. Include 20 g of agar if making YPD slants or plates.			
	3. Autoclave for 20 minutes on liquid cycle.			
	 Cool solution to ~55°C and add 100 ml of 20% Dextrose. Add 1.0 ml of 100 mg/ml Zeocin[™], if desired. 			
	Liquid medium without Zeocin TM can be stored at room temperature. Medium containing Zeocin TM should be stored at $+4^{\circ}$ C in the dark. YPD slants or plates are stored at $+4^{\circ}$ C. The shelf life of medium is several months. Medium containing Zeocin TM has a shelf life of one to two weeks.			

Recipes, continued

YPDS + Zeocin [™] Agar	Yeast Extract Peptone Dextrose Medium (1 liter)			
Agai	1% yeast extract			
	2% peptone			
	2% dextrose (D-glucose)			
	2% agar			
	100 µg/ml Zeocin [™]			
	1. Dissolve the following in 900 ml of water:			
	10 g yeast extract 182.2 g sorbitol			
	20 g of peptone			
	2. Add 20 g of agar.			
	3. Autoclave for 20 minutes on liquid cycle.			
	4. Add 100 ml of 20% Dextrose.			
	5. Cool solution to ~55°C and add 1.0 ml of 100 mg/ml Zeocin [™] .			
	Store YPDS slants or plates containing $\text{Zeocin}^{\text{TM}}$ at +4°C in the dark. The shelf life is one to two weeks.			
YNB + 0.5% Glucose	1.34 % YNB0.005% amino acids (if using YNB without amino acids)0.5% glucose			
	1. Dissolve 13.4 g YNB in 940 ml sterile water. (It may be necessary to heat the solution in order to dissolve the YNB).			
	2. Filter the solution. Autoclave 20 minutes on liquid cycle.			
	3. Add the following:			
	50 ml of a sterile 10% glucose solution			
	10 ml of a sterile 100X AA solution (if using YNB without amino acids)			
	Store at +4°C for approximately one year.			
Breaking Buffer	50 mM sodium phosphate, pH 7.4 1 mM PMSF (phenylmethylsulfonyl fluoride. You may use other protease inhibitors) 1 mM EDTA 5% glycerol			
	1. Prepare a stock solution of your desired protease inhibitors and store appropriately. Follow manufacturer's recommendations.			
	2. For 1 liter, dissolve the following in 900 ml deionized water:			
	6 g sodium phosphate (monobasic) 372 mg EDTA 50 ml glycerol			
	3. Use NaOH to adjust pH and bring up the volume to 1 liter. Store at $+4^{\circ}$ C.			
	4. Right before use, add the protease inhibitors.			

Zeocin

Zeocin[™] is a member of the bleomycin/phleomycin family of antibiotics isolated from Streptomyces. Antibiotics in this family are broad spectrum antibiotics that act as strong anti-bacterial and anti-tumor drugs. They show strong toxicity against bacteria, fungi (including yeast), plants, and mammalian cells (Baron et al., 1992; Drocourt et al., 1990; Mulsant et al., 1988; Perez et al., 1989).

The Zeocin[™] resistance protein has been isolated and characterized (Calmels *et al.*, 1991; Drocourt et al., 1990). This protein, the product of the Sh ble gene (Streptoalloteichus hindustanus bleomycin gene), is a 13.7 kDa protein that binds Zeocin[™] and inhibits its DNA strand cleavage activity. Expression of this protein in eukaryotic and prokaryotic hosts confers resistance to $\text{Zeocin}^{\text{TM}}$.

Molecular Weight, Formula, and Structure

The formula for ZeocinTM is $C_{60}H_{89}N_{21}O_{21}S_3$ and the molecular weight is 1,535. The diagram below shows the structure of $\text{Zeocin}^{\text{TM}}$.



Applications of Zeocin

Zeocin[™] is used for selection in mammalian cells (Mulsant et al., 1988); plants (Perez et al., 1989); yeast (Baron et al., 1992); and prokaryotes (Drocourt et al., 1990). Suggested concentrations of Zeocin^m for selection in *E. coli* and mammalian cells are listed below:

Organism	Zeocin [™] Concentration and Selective Medium	
E. coli	25-50 μg/ml in Low Salt LB medium* (see page 14 for recipe)	
Pichia	10-1000 µg/ml (varies with strain and medium)	
*Efficient selection requires that the concentration of NaCl be no more than 5 g/L (≤ 90 mM)		

Zeocin[™], continued

Handling Zeocin[™] High salt and acidity or basicity inactivate Zeocin[™]; therefore, we recommend that you reduce the salt in bacterial medium and adjust the pH to 7.5 to keep the drug active (see Low Salt LB Medium, page 14). Please note that the salt concentration and pH do not need to be adjusted when preparing tissue culture medium containing Zeocin[™]. Store Zeocin[™] at -20°C and thaw on ice before use. Zeocin[™] is light sensitive. Store drug, plates, and medium containing drug in the dark. Wear gloves, a laboratory coat, and safety glasses or goggles when handling solutions containing Zeocin[™]. Zeocin[™] is toxic. Do not ingest or inhale solutions containing the drug.

• Store tissue culture medium containing Zeocin[™] at +4°C in the dark. Medium containing Zeocin[™] is stable for 1-2 weeks.

Lithium Chloride Transformation Method

Introduction	This is a modified version of the procedure described for <i>S. cerevisiae</i> (Gietz and Schiestl, 1996). This protocol is provided as an alternative to transformation by electroporation. Transformation efficiency is between 10^2 to 10^3 cfu/µg linearized DNA.		
Preparation of Solutions	 Lithium acetate does not work with <i>Pichia pastoris</i>. Use only lithium chloride. 1 M LiCl in distilled, deionized water. Filter sterilize. Dilute as needed with sterile water. 50% polyethylene glycol (PEG-3350) in distilled, deionized water. Filter sterilize. Store in a tightly capped bottle. 2 mg/ml denatured, fragmented salmon sperm DNA in TE (10 mM Tris-HCl, pH 8.0, 		
	1.0 mM EDTA). Store at -20°C.		
Preparation of Cells	 Grow a 50 ml culture of <i>Pichia pastoris</i> in YPD at 30°C with shaking to an OD₆₀₀ of 0.8 to 1.0 (approximately 10⁸ cells/ml). 		
	 Harvest the cells and wash with 25 ml of sterile water. Centrifuge at 1500 x g for 10 minutes at room temperature. 		
	3. Decant the water and resuspend the cells in 1 ml of 100 mM LiCl.		
	4. Transfer the cell suspension to a 1.5 ml microcentrifuge tube.		
	5. Pellet the cells at maximum speed for 15 seconds and remove the LiCl with a pipet.		
	6. Resuspend the cells in 400 μl of 100 mM LiCl.		
	 Dispense 50 µl of the cell suspension into a 1.5 ml microcentrifuge tube for each transformation and use immediately. Do not store on ice or freeze at -20°C. 		
Transformation	 Boil a 1 ml sample of single-stranded DNA for five minutes, then quickly chill in ice water. Keep on ice. Note: It is not necessary nor desirable to boil the carrier DNA prior to each use. Store a small aliquot at -20°C and boil every 3-4 times the DNA is thawed. 		
	 Centrifuge the LiCl-cell solution from Step 7, above, and remove the LiCl with a ninet 		
	 3. For each transformation sample, add the following reagents IN THE ORDER GIVEN to the cells. PEG shields the cells from the detrimental effects of the high concentration of LiCl. 240 µl 50% PEG 36 µl 1 M LiCl 25 µl 2 mg/ml single-stranded DNA 		
	Plasmid DNA (5-10 µg) in 50 µl sterile water		
	4. Vortex each tube vigorously until the cell pellet is completely mixed (~ 1 minute).		
	5. Incubate the tube at 30° C for 30 minutes without shaking.		
	6. Heat shock in a water bath at 42°C for 20-25 minutes.		
	7. Centrifuge the tubes at 6000 to 8000 rpm and remove the transformation solution with a pipet.		
	8. Resuspend the pellet in 1 ml of YPD and incubate at 30°C with shaking.		
	 After 1 hour and 4 hours, plate 25 to 100 µl on YPD plates containing 100 µg/ml Zeocin[™]. Incubate the plates for 2-3 days at 30°C. Proceed to Analysis of <i>Pichia</i> Transformants, page 22. 		

Technical Service

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Product Qualification

Introduction

This section describes the criteria used to qualify the Pichia expression vectors

Vectors

Each vector is qualified by restriction enzyme digestion with specific restriction enzymes as listed below. Restriction digests must demonstrate the correct banding pattern when electrophoresed on an agarose gel.

Vector	Restriction Enzyme	Expected fragments (bp)
pGAPZ A	Apa I	2284
	Xba I	No sites
	ApaL 1	2214, 670
pGAPZ B	Apa I	No sites
	Xba I	2882
	ApaL I	2212, 670
pGAPZ C	Xba I	No sites
	SnaB I	2883
	ApaL I	2213, 670
pGAPZa A	Pst I	No sites
	Cla I	No sites
	ApaL I	2477, 670
pGAPZa B	Xba I	3151
	Pst I	3151
	ApaL I	2481, 670
pGAPZα	Xba I	3152
	Cla I	3152
	ApaL I	2482, 670

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