# The user manual: flavor Symmetry Embedded - GLoBES (FaSE-GLoBES)

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### I. OVERVIEW FASE-GLOBES

Flavor Symmetry Embedded (FaSE) is a supplemental tool for General Long Baseline Experiment Simulator (GLoBES) [1, 2] in order to analyse how a leptonic flavor symmetry model can be tested in neutrino oscillation experiments. FaSE is written in c/c++, and consist with two codes FaSE\_GLoBES.c and model-input.c. The user defines the model in to model-input.c, while FaSE\_GLoBES.c does not need to be touched.

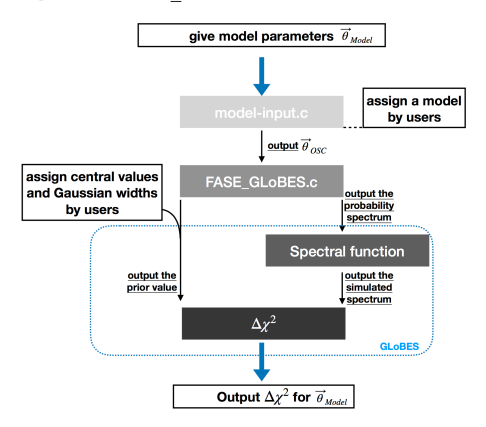


Figure 1: A sketch for the concept of FaSE-GLoBES. From top to down, this sketch shows three parts: 1. the parameter translation (model-input.c), 2. giving oscillation-parameter values (FaSE GLoBES.c), and 3. the  $\chi^2$ -value calculation (GLoBES).

Combining **GLoBES** with **Fase** (we call it '**FaSE-GLoBES**'), the user can analyse flavor symmetry models with the simulated experimental configuration. The concept of **FaSE-GLoBES** is shown in Fig. 1, in which three parts are shown: 1. **the parameter translation**, 2. **giving oscillation-parameter values**, and 3. **the**  $\chi^2$ -value calculation. The idea behind this flow chart in Fig. 1 is that given a set of model parameters  $\vec{\theta}_{Model}^{hyp}$  as a hypothese, the corresponding values in standard oscillation parameters  $\vec{\theta}_{OSC}$  are obtained

by a translation function, which is assigned by the user in **model-input.c**. And then, through **FaSE\_GLoBES.c** these oscillation-parameter values are passed into **GLoBES** library to simulate the event spectra so that the user can perform physics analysis with the newly-defined  $\chi^2$ .

API functions in **FaSE** are listed:

```
1. MODEL_init(N_{para}),
```

- 2. FASE\_glb\_probability\_matrix,
- 3. FASE\_glb\_set\_oscillation\_parameters,
- 4. FASE\_glb\_get\_oscillation\_parameters,
- 5. FASE\_prior\_OSC,
- 6. FASE\_prior\_model.

The first one is to initialise **FaSE** with the number of input parameters  $N_{para}$ . The next three functions need to be included to replace the default **GLoBES** probability engine. In the main code, the user needs to include the script as follows.

```
glbRegisterProbabilityEngine(6,
&FASE_glb_probability_matrix,
&FASE_glb_set_oscillation_parameters,
&FASE_glb_get_oscillation_parameters,
NULL);
```

This probability engine can work with oscillation or model parameters. It can be set by the user with the parameter PARA. If PARA=STAN (PARA=MODEL) the probability engine works with oscillation (model) parameters. The final two items on the API list are prior functions. Once the user gives the prior in oscillation (model) parameters, the user needs to call FASE\_prior\_OSC (FASE\_prior\_model) as follows.

```
glbRegisterPriorFunction(FASE_prior_OSC,NULL,NULL,NULL);
or
glbRegisterPriorFunction(FASE_prior_model,NULL,NULL,NULL);
```

We note that except for setting the probability engine and the prior function, the other parts

in the main code should follow with the GLoBES manual. Many simulation packages are provided on internet. Some of them are also available on **GLoBES** website [3], while the working group in the DUNE experiment also releases their neutrino flux information and detector descriptions in a AEDL file, provided in [4]. We summarise AEDL files for some experiments in Table. I, including their sources and references.

Table I: Summary of available AEDL files for some of most-discussed experiments.

exp.	source	ref.
T2K	T2K.glb on [3]	[5]
NOvA	NOvA.glb on [3]	[6]
T2HK	sys-T2HK.glb on [3]	[?]
DUNE	[4]	[4]
MOMENT	on FaSE website	[7]

In the following, we give more details about using **FASE-GLoBES**. In Sec. IA, we will give the instruction to download and install **GLoBES** and **FASE**, and we will introduce how to compile and run the code. More complicated is setting the model, and the way to do this will be described in Sec. IB. Finally, we will discuss how to set up the prior in Sec. IC.

### A. Download, compile and run

The user can download FaSE simply from https://github.com/tcwphy/FASE\_GLoBES.git directly. Or, the user can also download it:

git clone https://github.com/tcwphy/FASE\_GLoBES.git

As FaSE works based on GLoBES, the user should use the GLoBES Makefile, but include two binary files of FaSE. To do so, we include the script in the Makefile,

{makefile\_execution}: my\_program.o FASE\_GLoBES.o model-input.o
gcc my\_program.o FASE\_GLoBES.o model-input.o -o my\_executable \$(LDFLAGS)
\$(local\_LDFLAGS)

where {compile\_execusion} is the makefile commend for the program my\_program.o and the execution my\_executable is the output. After giving this script in the GLoBES

Makefile, the user needs to makefile the program my\_program on the terminal.

makefile {makefile\_execution}

And, the user can execute the program by typing the following commend on the terminal.

./my\_executable

The user can also compile the program without the **GLoBES** Makefile with the following script.

gcc my\_program.o FASE\_GLoBES.o model-input.o -lglobes -LGLB\_DIR/lib/ -o
my\_executable<sup>1</sup>

And, the execution is also in the same way: ./my\_executable.

Finally we need to initialize **FaSE** by giving  $N_{para}$  which is the number of model parameters. To do so, we need to include the following script in the main code.

 $\texttt{MODEL\_init}(N_{para});$ 

We note that  $N_{para}$  should not be larger than the number of standard oscillation parameters, 6.

# B. Model setting

The function MtoS can do the translation from model parameters  $\vec{\theta}_{Model}$  to oscillation parameters  $\vec{\theta}_{OSC}$ . After the user gives the array  $\vec{\theta}_{Model}$  to the function MtoS, the output is the corresponding oscillation parameter  $\vec{\theta}_{OSC}$ , of which components are  $\theta_{12}$ ,  $\theta_{13}$ ,  $\theta_{23}$ ,  $\delta$ ,  $\Delta m_{21}^2$ , and  $\Delta m_{31}^2$ . For the first four components, values are given in the unit of **rad**, while the other two are in **eV**<sup>2</sup>. These values will be passed into **FaSE\_GLoBES.c** to simulate experimental spectra and compute prior values.

There are three methods to translate from  $\vec{\theta}_{Model}$  to  $\vec{\theta}_{OSC}$  in **FaSE-GLoBES** as follows.

1. Assign the correlation between the standard oscillation and model parameter sets. In the way of directly giving the relation between oscillation and model parameter sets,

<sup>&</sup>lt;sup>1</sup> We note that only the gcc compiler has been tested. Users can also try other compilers.

the user needs to provide

$$\vec{\theta}_{Model} = \vec{f}(\vec{\theta}_{OSC}),\tag{1}$$

in the function MtoS.

2. Give the mixing matrix in model parameter U. When the user gives the mixing matrix U in model parameters, the corresponding mixing angles can be obtained through relations,

$$\tan \theta_{12} = \left| \frac{U_{e2}}{U_{e1}} \right|, \sin \theta_{13} = |U_{e3}|, \tan \theta_{23} = \left| \frac{U_{\mu 3}}{U_{\tau 3}} \right|.$$
(2)

After getting these mixing angles, we can easily get the Dirac CP phase  $\delta$  with the Jarlskog invariant  $J_{CP}$ ,

$$J_{CP} = c_{12} s_{12} c_{23} s_{23} c_{13}^2 s_{13} \sin \delta, \tag{3}$$

where  $c_{ij}$  and  $s_{ij}$  are  $\cos \theta_{ij}$  and  $\sin \theta_{ij}$ , respectively. In MtoS, the user can pass the mixing matrix U to the function STAN\_OSC\_U to obtain the corresponding mixing angles and the CP phase.

3. Define the mass matrix in model parameters. The oscillation parameters can also be obtained in the way based on

$$U^{\dagger} \mathcal{M} \mathcal{M}^{\dagger} U = \mathbf{M}^2$$
, where  $\mathbf{M}_{\alpha\beta}^2 = m_{\alpha}^2 \delta_{\alpha\beta}$ , (4)

where  $\mathcal{M}$  (**M**) is the neutrino mass matrix in the flavour (mass) state. The matrix  $\mathcal{M}$  is given by the user with model parameters  $\vec{\theta}_{Model}$ . The mixing matrix U can be used to get mixing angles and the CP phase, as Eqs. (2) and (3). The difference between any two diagonal elements of  $\mathbf{M}$  ( $\mathbf{M}_{ii} - \mathbf{M}_{jj} = m_i^2 - m_j^2$ ) is the mass-squared difference ( $\Delta m_{ij}^2$ ). This diagnolisation in Eq. (4) can be done by the function STAN\_OSC, which needs to be called in MtoS with outputs of the vector  $\vec{\theta}_{OSC}$ . We note that with non-diagonal mass matrix for charged leptons, this translation method is not suggested.

More details about how to assign a model will be introduced in Sec. II A.

### C. Prior setting

Given a set of values for model parameters, FaSE\_GLoBES.c will obtain the corresponding oscillation-parameter values from model-input.c, and will pass these values

to simulate event spectra and to compute the prior value. Two gaussian prior functions are provided in FaSE: FASE\_prior\_OSC and FASE\_prior\_model. These two functions are for different purposes. If the user gives the prior in oscillation (model) parameters, the user should register FASE\_prior\_OSC (FASE\_prior\_model) with the GLoBES function glbRegisterPriorFunction, as we introduced in the beginning of this section. The user also needs to assign the parameter PARA=STAN (PARA=Model), when the user prefers to give the prior in oscillation (model) parameters. The Gaussian prior is

$$\chi_{prior}^2 = \sum_i \frac{(\theta_i - \theta_i^c)^2}{\sigma_i^2},\tag{5}$$

where  $\theta_i$  is one parameter of the hypothesis  $\vec{\theta}^{hyp}$ ,  $\theta_i^c$  ( $\sigma_i$ ) is the central value (Gaussian width) of the prior for  $\theta_i$ . We note that  $\vec{\theta}^{hyp}$  can be either model ( $\vec{\theta}_{Model}$ ) or oscillation parameters ( $\vec{\theta}_{OSC}$ ). The values of  $\theta_i^c$  and  $\sigma_i$  need to be given by the user through three arrays: Central\_prior, UPPER\_prior, and LOWER\_prior, in which there are six components. To treat asymmetry of width for upper ( $\theta_i > \theta_i^c$ ) and lower ( $\theta_i < \theta_i^c$ ) Gaussian widths, we give values in two arrays UPPER\_prior and LOWER\_prior, respectively. Once the user does not want to include any priors, two arrays UPPER\_prior and LOWER\_prior need to be 0. If the user gives the prior in model parameters, the order of each component follows with the setup of input for the probability engine. If the user gives the prior in oscillation parameters, the six components in order are  $\theta_{12}$ ,  $\theta_{13}$ ,  $\theta_{23}$ ,  $\delta$ ,  $\Delta m_{21}^2$ , and  $\Delta m_{31}^2$ . The first four parameter are in rad, and the final two are in eV<sup>2</sup>.

Finally, some restrictions are imposed by the studied flavor symmetry model. We set up these restrictions in the function model\_restriction in model\_input.c. In the function model\_restriction, the user needs to return 1 once the restriction is broken. For example, if the normal ordering is imposed, we give "if (DMS31<0) { return 1;} "in model\_restriction, where DMS31 is the variable for  $\Delta m_{31}^2$ . If there is no any restrictions, we simply return 0 in model\_restriction as follows.

double model\_restriction(double model []){ return 0;}

# D. The definition of $\chi^2$ function

The user can use **FaSE-GLoBES** to constrain model parameters. Suppose we have the measurement  $\vec{x}$  and the likelihood function  $L(\vec{\theta}^{hyp}) = P(\vec{x}|\vec{\theta}^{hyp})$  for a set of parameters  $\vec{\theta}^{hyp} = (\theta_1, ..., \theta_N)$ , where  $P(\vec{x}|\vec{\theta}^{hyp})$  is the probability function for data  $\vec{x}$  in favour of the hypothesis  $\vec{\theta}^{hyp}$ . The constraint of model parameters can be obtained with the statistis parameter  $\chi^2 \equiv \ln L(\vec{\theta}^{hyp})$ . The expression  $\chi^2$  is used as the default **GLoBES** setting. In more detail, the  $\chi^2$  function, following the Poisson distribution, is constructed based on a log-likelihood ratio,

$$\chi^{2}(\vec{\theta}^{hyp}, \xi_{s}, \xi_{b}) = 2\sum_{i} \left( \eta_{i}(\vec{\theta}^{hyp}, \xi_{s}, \xi_{b}) - n_{i} + n_{i} \ln \frac{n_{i}}{\eta_{i}(\vec{\theta}^{hyp}, \xi_{s}, \xi_{b})} \right) + p(\xi_{s}, \sigma_{s}) + p(\xi_{b}, \sigma_{b}) + \chi^{2}_{prior},$$

$$(6)$$

where i runs over the number of bins,  $\eta_i(\vec{\theta}, \xi_s, \xi_b)$  is the assumed event rate in the ith bin and  $E_i$  is the central value in this energy bin. The vector  $\vec{\theta}$  consists of model or oscillation parameters. The parameters  $\xi_s$  and  $\xi_b$  are introduced to account for the systematic uncertainties in the normalisation for the signal (subscript  $_s$ ) and background (subscript  $_b$ ) components of the event rate, and are allowed to vary in the fit as nuisance parameters. For a given set of parameters  $\vec{\theta}^{hyp}$ , the event rate in the ith energy bin is calculated as

$$\eta_i(\vec{\theta}^{hyp}, \xi_s, \xi_b) = (1 + \xi_s) \times s_i + (1 + \xi_b) \times b_i, \tag{7}$$

where  $s_i$  and  $b_i$  are the expected number of signal and background events in *i*th energy bin, respectively. The nuisance parameters are constrained by the Gaussian prior  $p(\xi, \sigma) = \xi^2/\sigma^2$  with corresponding uncertainties  $\sigma_s$  and  $\sigma_b$  for the signal and background, respectively. Finally,  $\chi^2_{prior}$  is a set of Gaussian priors for hypothesis, and is expressed as Eq. (5). After doing all minimisations, the user obtains the  $\chi^2$  value for a specific hypothesis  $\vec{\theta}^{hyp}$ ,  $\chi^2(\vec{\theta}^{hyp})$ .

Based on the  $\chi^2$  function Eq. (6), we can study how model parameters can be constrained and whether a flavor-symmetry neutrino model is excluded by simulated experiments. In the following we will demonstrate how it works.

### Applications

The user of **FaSE-GLoBES** is able to study how model parameters can be constrained by the simulated experiments. To do so, the user needs to simulate the true event spectrum  $n_i$  with a set of model  $(\vec{\theta}_{Model}^{true})$  or oscillation parameters  $(\vec{\theta}_{OSC}^{true})$ , *i.e.* set up  $n_i(\vec{\theta}_{Model}^{true})$  or  $n_i(\vec{\theta}_{OSC}^{true})$ . The hypothesis  $\vec{\theta}_{Model}^{hyp}$  predicts the tested event spectrum  $\eta_i(\vec{\theta}_{Model}^{hyp}, \xi_s, \xi_b)$ . With the default settings for  $\chi^2$  function as Eq. (5) in **FaSE-GLoBES**, the user computes the statistical quantity,

$$\chi^2(\vec{\theta}_{Model}^{hyp})$$
, with  $n_i(\vec{\theta}_{Model}^{true})$  or  $n_i(\vec{\theta}_{OSC}^{true})$ . (8)

We note that the minimum of  $\chi^2$  in the whole parameter space  $(\chi^2_{min.})$  may not be 0. Therefore, to get the precision of model parameters, the user should use the value  $\Delta\chi^2(\vec{\theta}^{hyp}_{Model}) \equiv \chi^2(\vec{\theta}^{hyp}_{Model}) - \chi^2_{min.}$ , instead of  $\chi^2(\vec{\theta}^{hyp}_{Model})$  itself. By varying different hypotheses  $\vec{\theta}^{hyp}_{Model}$ , we will obtain the allowed region of model parameters with the statistical quantity  $\Delta\chi^2(\vec{\theta}^{hyp}_{Model})$ .

The user can also study how well a flavor symmetry model explains the computed data, or predict whether the simulated experiment can exclude this model or not. In other words, the user studies the minimum of  $\chi^2$  for the flavor symmetry model  $\vec{\theta}_{Model}$  as a hypothesis, by assuming different true oscillation values, *i.e.* different  $\vec{\theta}_{OSC}^{true}$ . To do so, one can compute the same statistical quantity in Eq. (8), while the true spectrum is varied with different true values  $\vec{\theta}_{OSC}^{true}$ . All model parameters are allowed to be varied with the user-defined prior. Finally, the user might adopt Wilk's theorem to interpret results [8]. When we compare nested models, the  $\Delta\chi^2$  test statistics is a random variable asymptotically distributed as a  $\chi^2$ -distribution with the number of degrees of freedom, which is equal to the difference in the number of free model parameters.

In Secs. II D and II E, we will present examples to demonstrate how the user can do with **FaSE-GLoBES** to constrain the model parameter and to exclude a model by the simulated experiment configuration, respectively.

# II. EXAMPLES

In this manual, we now take the tri-direct littlest seesaw (TDLS) [10–12] and a  $S_4$  modular flavor symmetry model [13] as examples, and simulate the event spectra for Deep Under-

Table II: A summary of the relation between oscillation parameters and TDLS model parameters [9]. Two requirements are imposed by TDLS: the smallest mass state  $m_1 = 0$  and the normal mass ordering. The sign of  $\sin \delta$  depends on the sign of  $x \cos \psi$ : "+" ("-") is for  $x \cos \psi > 0$  (< 0).

model parameters	$x, \eta, r, m_a$			
	$y = \frac{5x^2 + 2x + 2}{2(x^2 + x + 1)}(m_a + e^{i\eta}m_s),$			
	$z = -\frac{\sqrt{5x^2 + 2x + 2}}{2(x^2 + x + 1)} \left[ (x + 2)m_a - x(2x + 1)e^{i\eta}m_s \right],$			
combinations of model parameters	$w = \frac{1}{2(x^2+x+1)} \left[ (x+2)^2 m_a + x^2 (2x+1)^2 e^{i\eta} m_s \right],$			
	$\sin \psi = \frac{\Im(y^*z + wz^*)}{ y^*z + wz^* },  \cos \psi = \frac{\Re(y^*z + wz^*)}{ y^*z + wz^* }.$			
	$\sin 2\theta = \frac{2 y^*z + wz^* }{\sqrt{( w ^2 -  y ^2)^2 + 4 y^*z + wz^* ^2}},$			
	$\cos 2\theta = \frac{ w ^2 -  y ^2}{\sqrt{( w ^2 -  y ^2)^2 + 4 y^*z + wz^* ^2}}.$			
	$\Delta m_{21}^2 = m_2^2 = \frac{1}{2} \left[  y ^2 +  w ^2 + 2 z ^2 - \frac{ w ^2 -  y ^2}{\cos \theta} \right],$			
	$\Delta m_{31}^2 = m_3^2 = \frac{1}{2} \left[  y ^2 +  w ^2 + 2 z ^2 + \frac{ w ^2 -  y ^2}{\cos \theta} \right],$			
oscillation parameters	$\sin^2 \theta_{12} = 1 - \frac{3x^2}{3x^2 + 2(x^2 + x + 1)\cos^2 \theta},$			
	$\sin^2 \theta_{13} = \frac{2(x^2 + x + 1)\sin^2 \theta}{5x^2 + 2x + 2},$			
	$\sin^2 \theta_{23} = \frac{1}{2} + \frac{x\sqrt{3(5x^2 + 2x + 2)}\sin 2\theta \sin \psi}{2[3x^2 + 2(x^2 + x + 1)\cos^2 \theta]},$			
	$\cos \delta = \frac{\cot 2\theta_{23} \left[ 3x^2 - \left( 4x^2 + x + 1 \right) \cos^2 \theta_{13} \right]}{\sqrt{3}  x  \sin \theta_{13} \sqrt{(5x^2 + 2x + 2) \cos^2 \theta_{13} - 3x^2}},$			
	$\sin \delta = \pm \csc 2\theta_{23} \sqrt{1 + \frac{(x^2 + x + 1)^2 \cot^2 \theta_{13} \cos^2 2\theta_{23}}{3x^2 [3x^2 \tan^2 \theta_{13} - 2(x^2 + x + 1)]}}.$			

ground Neutrino Experiment (DUNE) [14] and the MuOn-decay MEdium baseline NeuTrino beam experiment (MOMENT) [7] by GLoBES. In TDLS model, the atmospheric and solar flavon vacuum alignments are  $\langle \phi_{\rm atm} \rangle \propto (1, \omega^2, \omega)^T$  and  $\langle \phi_{\rm sol} \rangle \propto (1, x, x)^T$ , where  $\omega$  stands for a cube root of unity and the parameter x is real because of the imposed CP symmetry. Under this model, the light left-handed Majorana neutrino mass matrix in the flavor basis is given by

$$m_{\nu} = m_a \begin{pmatrix} 1 & \omega & \omega^2 \\ \omega & \omega^2 & 1 \\ \omega^2 & 1 & \omega \end{pmatrix} + e^{i\eta} m_s \begin{pmatrix} 1 & x & x \\ x & x^2 & x^2 \\ x & x^2 & x^2 \end{pmatrix}, \tag{9}$$

where x,  $\eta$ ,  $m_a$ , and the ratio  $r \equiv m_s/m_a$  are four parameters and will be constrained by experimental data. And based on Eq. (9), the relation between the model and oscillation parameters are predicted as Table. II. We note that from Eq. (9),  $m_1 = 0$  and the normal

mass ordering are imposed, and will need to be imposed in **FaSE-GLoBES**. Therefore, the restrictions in this model are  $m_a > 0$  and r > 0.

In the  $S_4$  modular model, the neutrino mass is predicted:

$$m_{\nu} = (\mu_{1}\hat{c}_{R}^{2} + \mu_{2}\hat{s}_{R}^{*2}) \begin{pmatrix} 1 & -2\omega^{2} & -2\omega \\ -2\omega^{2} & 4\omega & 4 \\ -2\omega & 4 & 4\omega^{2} \end{pmatrix} + (\mu_{1}\hat{s}_{R}^{2} + \mu_{2}\hat{c}_{R}^{*2}) \begin{pmatrix} 0 & 0 & 0 \\ 0 & 1 & -1 \\ 0 & -1 & 1 \end{pmatrix} + (\mu_{1}\hat{c}_{R}\hat{s}_{R} - \mu_{2}\hat{c}_{R}^{*}\hat{s}_{R}^{*}) \begin{pmatrix} 0 & -1 & 1 \\ -1 & 4\omega^{2} & 2i\sqrt{3} \\ 1 & 2i\sqrt{3} & -4\omega \end{pmatrix},$$
(10)

where  $\hat{c}_R$  and  $\hat{s}_R$  are  $\cos \theta_R \times e^{i\alpha_2}$  and  $\sin \theta_R \times e^{i\alpha_3}$ , respectively. As  $\omega$  is fixed at  $-\frac{1}{2} + i\frac{\sqrt{3}}{2}$ , this model has 5 model parameters:  $\mu_1$ ,  $\mu_2$ ,  $\theta_R$ ,  $\alpha_2$ ,  $\alpha_3$ .

In the following, all examples are mainly presented for TDLS, except for Sec. II E. However, we remind the user that though we use these two models as examples, **FaSE-GLoBES** should not be limited in any flavor model. We therefore in the **FaSE** distribution, show the warped flavor symmetry (WSF) [15], trimaximal mixing sum rule (TM1) [16] as examples, which will not be further introduced in this manual. We summarise our examples for **FaSE-GLoBES** and references for models themselves in Tabl III. And, in the rest of this section, we demonstrate the **example 1** and **example 4**.

Table III: Summary of available AEDL files for some of most-discussed experiments.

example	model	application	reference
example1	TDLS	parameter constraint	[10-12]
example2	WSF	parameter constraint	[15]
example3	TM1 sum rule	model testing	[16]
example4	$S_4$ modular	model testing	[13]

### A. Setup for a model

If the user prefers to assign the model with the mass matrix, the user will need to define the complex matrix Mass\_Matrix with Eq. (9) in the function MtoS. The user can therefore assign the model as follows.

```
int MtoS(double OSC_PARAMS[6], double M_para[])
{
double x=M_para[0];
double eta=M_para[1];
double r=M_para[2];
double ma=M_para[3];
double ms=ma*r;
          double complex Mass_Matrix[] = {ma+ms*(cos(eta) + I*sin(eta)),
ma*(cos(6.6666e-1*M_PI) + I*sin(6.6666e-1*M_PI))+x*ms*(cos(eta) + I*ma*(cos(6.6666e-1*M_PI))+x*ms*(cos(eta) + I*ma*(cos(6.6666e-1*M_PI))+x*ms*(cos(eta) + I*ma*(eta))+x*ms*(cos(eta) + I*ma*(eta))+x*ms*(eta))+x*ms*(eta) + I*ma*(eta))+x*ms*(eta) + I*ma*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))
I*sin(eta)), ma*(cos(6.6666e-1*M_PI) + I*sin(6.6666e-1*M_PI))*(cos(6.6666e-1*M_PI))
+ I*sin(6.666e-1*M_PI))+x*ms*(cos(eta) + I*sin(eta)),
ma*(cos(6.6666e-1*M_PI) + I*sin(6.6666e-1*M_PI))+x*ms*(cos(eta) + I*ma*(cos(6.6666e-1*M_PI))+x*ms*(cos(eta) + I*ma*(cos(6.6666e-1*M_PI))+x*ms*(cos(eta) + I*ma*(eta))+x*ms*(cos(eta) + I*ma*(eta))+x*ms*(eta) + I*ma*(eta))+x*ms*(eta) + I*ma*(eta))+x*ms*(eta) + I*ma*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta))+x*ms*(eta)
I*sin(eta)), ma*(cos(6.6666e-1*M_PI) + I*sin(6.6666e-1*M_PI))*(cos(6.6666e-1*M_PI))
+ I*sin(6.6666e-1*M_PI))+x*x*ms*(cos(eta) + I*sin(eta)),
ma+x*x*ms*(cos(eta) + I*sin(eta)), ma*(cos(6.6666e-1*M_PI) +
I*sin(6.6666e-1*M_PI))*(cos(6.6666e-1*M_PI) + I*sin(6.6666e-1*M_PI))+x*ms*(cos(eta))
+ I*sin(eta)), ma+x*x*ms*(cos(eta) + I*sin(eta)), ma*(cos(6.6666e-1*M_PI) +
I*sin(6.6666e-1*M_PI))+x*x*ms*(cos(eta) + I*sin(eta));
          STAN_OSC(Mass_Matrix,OSC_PARAMS);
          return 0;
}
           The user can also provide the relation between model and oscillation parameters Eq. (1),
such as Table. II. The user, who prefers to do so, can assign a model as follows.
           double MtoS(double osc_para[6], double M_para[])
{
/*example: tri-direct*/
double x=M_para[0];
double eta=M_para[1];
```

```
double r=M_para[2];
double ma=M_para[3];
osc_para[0]=TDth12(x,eta,r, ma);
osc_para[1]=TDth13(x,eta,r, ma);
osc_para[2]=TDth23(x,eta,r, ma);
osc_para[3]=TDdCP(x,eta,r, ma);
osc_para[4]=TDdm21(x,eta,r, ma);
osc_para[5]=TDdm31(x,eta,r, ma);
return 0;
}
   The functions TDth12, TDth13, TDth23, TDdCP, TDdm21, and TDdm31 are included for
obtaining the value of \theta_{12}, \theta_{13}, \theta_{23}, \delta, \Delta m_{21}^2, and \Delta m_{31}^2 according to Table II, respectively.
   The restrictions m_a > 0 and r > 0 need to be setup in model_restriction as follows.
double model_restriction(double model [])
{
double x=model[0];
double eta=model[1];
double r=model[2];
double ma=model[3];
if(ma<0) {return 1;}</pre>
if(r<0) {return 1;}</pre>
return 0;
}
```

### B. Initialise the code

In the beginning of main code, we need to initialise **GLoBES** and **FaSE**, and to include the AEDL file for MOMENT as follows.

```
glbInit(argv[0]);
glbInitExperiment("exp/MOMENT_FIX_FLUX_150KM_addATM_NC.glb",&glb_experiment_list[0],
&glb_num_of_exps);
MODEL_init(4);
Then we register the probability engine.
glb_init_probability_engine();
glbRegisterProbabilityEngine(6,
&FASE_glb_probability_matrix,
&FASE_glb_set_oscillation_parameters,
&FASE_glb_get_oscillation_parameters,
NULL);
In the following we define the model-parameter values (x, \eta, r, \text{ and } m_a), and set the true
value in model parameters true_values to simulated the true event spectrum. Here we
note that for setting up the true spectrum with model (oscillation) parameters, the user
needs to make PARA equal to MODEL (STAN).
float x_true,eta_true,r_true,ma_true;
x_true=-3.65029; eta_true=1.13067*M_PI; r_true=0.511325;
ma_true=3.71199e-3;
glb_params true_values = glbAllocParams();
glbDefineParams(true_values,x_true,eta_true,r_true,ma_true,0,0);
glbSetDensityParams(true_values,1.0,GLB_ALL);
PARA=MODEL;
glbSetOscillationParameters(true_values);
glbSetRates();
```

### C. Prior setup

To set up the prior, we need to register the prior function. As mentioned, when we use the prior in oscillation and model parameters, we register FASE\_prior\_OSC and FASE\_prior\_model, respectively. Here we present an example setting up priors in oscillation parameters according to NUFIT4.0 Table IV.

						. ,
Parameter	$\theta_{12}/^{\circ}$	$\theta_{13}/^{\circ}$	$\theta_{23}/^{\circ}$	$\delta/^{\circ}$	$\Delta m_{21}^2 / 10^{-5} \text{eV}^2$	$\Delta m_{31}^2 / 10^{-3} \text{eV}^2$
best fit	33.82	8.61	49.6	215	7.39	2.525
$3\sigma$ Range	31.61 - 36.27	8.22 - 8.99	40.3 - 52.4	125 - 392	6.79 - 8.01	2.47 - 2.625

Table IV: The best fit and  $3\sigma$  uncertainty, in the results of NuFit4.0 [17].

We therefore register FASE\_prior\_OSC for the prior function as follows.

```
glbRegisterPriorFunction(FASE_prior_OSC, NULL, NULL, NULL);
```

And, then we give the value for the central values, and upper and lower Gaussian widths as follows.

```
UPPER_prior[0]=36.27*degree; LOWER_prior[0]=31.61*degree;
Central_prior[0]=33.82*degree;
UPPER_prior[1]=8.99*degree; LOWER_prior[1]=8.22*degree;
Central_prior[1]=8.61*degree;
UPPER_prior[2]=52.4*degree; LOWER_prior[2]=40.3*degree;
Central_prior[2]=49.6*degree;
UPPER_prior[3]=392*degree; LOWER_prior[3]=125*degree;
Central_prior[3]=215*degree;
UPPER_prior[3]=215*degree;
UPPER_prior[4]=8.01e-5; LOWER_prior[4]=6.79e-5; Central_prior[4]=7.39e-5;
UPPER_prior[5]=2.625e-3; LOWER_prior[5]=2.427e-3; Central_prior[5]=2.525e-3;
int i;
for (i=0;i<6;i++) {UPPER_prior[i]=fabs(UPPER_prior[i]-Central_prior[i])/3;}
LOWER_prior[i]=fabs(LOWER_prior[i]-Central_prior[i])/3;}
for (i=0; i<6; i++) glbSetOscParams(centers,0,i);
glbSetDensityParams(centers,1.0,GLB_ALL); glbCopyParams(centers,input_errors);</pre>
```

glbSetCentralValues(centers); glbSetInputErrors(input\_errors); The parameter degree is defined as  $\pi/180$ .

### D. Constraint of model parameters

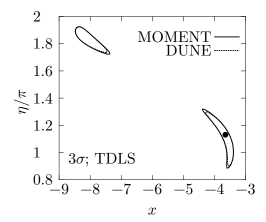


Figure 2: Example for using **FaSE-GLoBES** to obtain the constraints of model parameters for tri-direct littlest seesaw, with simulated DUNE and MOMENT data. The result is assumed the normal ordering. And, the dot denotes the model prediction with with NuFit4.0 results.

We show an example for how the model parameter can be constrained by MOMENT experiment, assuming the true model values x = -3.65,  $\eta = 1.13\pi$ , r = 0.511, and  $m_a = 3.71$  meV. To get the constraint of x and  $\eta$ , we need to compute the minimum of  $\chi^2$  value as the following script in the main code.

```
glbSetProjection(free);
float res0=glbChiNP(true_values,NULL,GLB_ALL);
```

The **GLoBES** projector free allows all model parameters to vary, in order to find the value of res0, which is  $\chi^2_{min}$ . Then, we set two loops varying different values of x and  $\eta$  in  $\vec{\theta}^{hyp}_{Model}$  to get  $\Delta\chi^2$  values for different hypotheses. We note here there is degenerate solutions where  $x \sim -8$ ,  $\eta \sim 1.8\pi$ ,  $r \sim 0.1$ , and  $m_a \sim 3.8$  meV. To find this degenerate solutions we set the other entry with r = 0.

float x,eta,r,ma,dx,deta;

```
float lower_x,upper_x,lower_eta,upper_eta;
FILE* File=fopen("data/constraint_x_eta.dat", "w");
lower_x=-9; upper_x=-3; lower_eta=0.8*M_PI; upper_eta=2*M_PI;
dx=(upper_x-lower_x)/100; deta=(upper_eta-lower_eta)/100;
glbSetProjection(projection);
for (x=lower_x;x<=upper_x;x=x+dx){
  for (eta=lower_eta;eta<=upper_eta;eta=eta+deta){
    glbSetOscParams(test_values,x,0); glbSetOscParams(test_values,eta,1);
    glbSetOscParams(test_values,r_true,2);
  float res=glbChiNP(test_values,NULL,GLB_ALL);
    glbSetOscParams(test_values,0.,2);
  float res2=glbChiNP(test_values,NULL,GLB_ALL);
    if(res2<res) {res=res2;}
    fprintf(File,"%f %f %f \n",x,eta/M_PI,res-res0);
} fprintf(File,"\n");}</pre>
```

The **GLoBES** projector projection fixes values of x and  $\eta$ , but allows the other parameters to vary. The output (data/contraint\_x\_eta.dat) is a set of  $\Delta \chi^2(\vec{\theta}^{hyp})$  (the third column) values with the changed x (the first column),  $\eta$  (the second column). We use these data, and obtain the predicted  $1\sigma$ ,  $2\sigma$ , and  $3\sigma$  constraints of x and  $\eta$  by MOMENT shown in Fig. 2. To get the result for DUNE, the user adopt the same way, but use different simulation package. The code for this example is provided as  $\mathbf{x}_{e}$  at  $\mathbf{x}_{e}$  at  $\mathbf{x}_{e}$  the FaSE distribution.

### E. Model testing

The user can also study how well a flavor symmetry model explains the computed data as predicting how the simulated experiment can exclude this model. As an example, we present testing a  $S_4$  modular symmetry model, assuming various true values of  $\theta_{23}$  and  $\delta$ . The code for this example is provided as **th23\_delta.c** in the **example4** of **FaSE** distribution. We compute the minimal  $\chi^2$  value for this  $S_4$  modular model with all model parameters, varied with the prior in Eq. (5) according to NuFit4.0 results. The studied

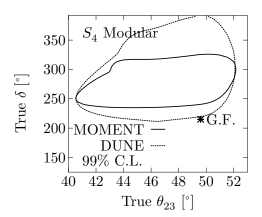


Figure 3: Example for using **FaSE-GLoBES** to obtain the 2-D exclusion contour at 99% C.L. on the plane of true values of  $\theta_{23}$  and  $\delta$  for a  $S_4$  modular symmetry model with DUNE and MOMENT. The result is assumed the normal ordering. And, the dot (star) denotes the model prediction with NuFit4.0 results (best fit of NuFit4.0).

statistics function is exactly Eq. (6), but the true event rate  $n_i$  is predicted by a set of oscillation parameters, which will be varied in the code. And all four model parameter can be varied with the prior given by Eq. (5). Except for  $\theta_{23}$  and  $\delta$ , which are varied between  $40^{\circ} < \theta_{23} < 52^{\circ}$  and  $125^{\circ} < \delta < 392^{\circ}$ , we fix the other oscillation parameters at the current best fit result Table IV. Finally, in the code we adopt Wilk's theorem [8]: textttgsl\_cdf\_chisq\_Qinv(gsl\_cdf\_chisq\_Q(fabs(res),dof),1).

```
float th23,dCP,dth23,ddCP,lower_th23,upper_th23,lower_dCP,upper_dCP;
FILE* File=fopen("data/S4Modular_th23_dCP(DUNE)_v3.dat", "w");
lower_th23=40; upper_th23=52.5; lower_dCP=125; upper_dCP=395;
dth23=.5; ddCP=5.;

int dof=1;
glbSetProjection(free);
for (th23=lower_th23;th23<=upper_th23;th23=th23+dth23){
    for (dCP=lower_dCP;dCP<=upper_dCP;dCP=dCP+ddCP){
        glbSetOscParams(true_values,th23*degree,GLB_THETA_23);
    }
}</pre>
```

```
glbSetOscParams(true_values,dCP*degree,GLB_DELTA_CP);
PARA=STAN;
glbSetOscillationParameters(true_values);
glbSetRates();
PARA=MODEL;
glbDefineParams(test_values,mu1,mu2,theta_R,alpha1,alpha2,0);
float res=glbChiNP(test_values,out,GLB_ALL);
for(double alpha2_test=25; alpha2_test<400; alpha2_test=alpha2_test+300){</pre>
glbSetOscParams(test_values,alpha2_test*degree,4);
for(double alpha1_test=60; alpha1_test<400; alpha1_test=alpha1_test+300){</pre>
glbSetOscParams(test_values,alpha1_test*degree,3);
glbSetOscParams(test_values,40*degree,2);
float res2=glbChiNP(test_values,out2,GLB_ALL);
if(res>res2) res=res2; glbCopyParams(out2,out);
glbSetOscParams(test_values, 20*degree, 2);
res2=glbChiNP(test_values,out2,GLB_ALL);
if(res>res2) res=res2; glbCopyParams(out2,out);
}}
 fprintf(File,"%f %f %f\n",th23,dCP,\
 gsl_cdf_chisq_Qinv(gsl_cdf_chisq_Q(fabs(res),dof),1));
 } fprintf(File,"\n");}
```

We note that hoops for variables alpha2\_text, alpha1\_test, and switch the entry value of  $\theta_R$  between 20° and 40° is for finding the degenerate solutions, according to Ref. [13]. The output data/S4Modular\_th23\_dCP\_test.dat contains true values of  $\theta_{23}$  (the first column) and  $\delta$  (the second column) with  $\chi^2_{min}$  for this  $S_4$  modular model (the third column). With these data, we then have the results shown in Fig. 3. To get the result for DUNE, the user

adopt the same way, but use different simulation package.

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