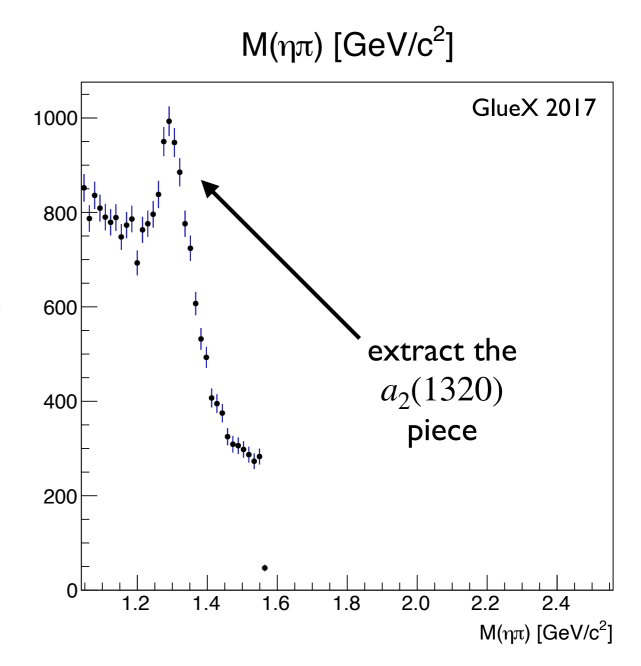
Using Amp Tools to Extract the $a_2(1320)$ Yield from GlueX 2017 Data

GlueX Software Tutorial May 23, 2022

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Example Fit Goal: Extract a_2 from $\eta \pi^0$

- Use data sample from previous exercise
- The kinematics of $\eta \pi^0$ system (angular distributions) are given by the Z_ℓ^m functions discussed in the morning session
- Dynamical assumptions about how the amplitude depends on $M(\eta\pi^0)$
 - D-waves: a complex-valued Breit-Wigner with parameters consistent with the $a_2(1320)$
 - certain m-projections/reflectivities excluded based on tensor meson dominance (TMD) model
 - S-waves: fixed complex number in four coarse bins of $M(\eta \pi^0)$ (a piecewise complex function)
- Goal: extract the total efficiency corrected yield of $a_2(1320)$ (the coherent sum of all D-waves)





Required Code: AmpTools

- AmpTools: library that is distributed independently of GlueX code base
 - available through GitHub: <u>github.com/mashephe/AmpTools</u>
 - installed as external package in GlueX environment
 - (include version.xml lines here)
- The AmpTools library (not an executable) that provides a general interface for doing fits
 - fully functional example of how to use AmpTools is also provided in <u>Tutorials/Dalitz</u>
 - <u>copyDalitz.py</u> is provided to generate your own project based on the Dalitz tutorial (not needed for a typical GlueX user)
 - GlueX has a set of libraries and executables that rely on the core AmpTools package
 - halld_sim/src/libraries/AMPTOOLS_...
 - halld_sim/src/programs/AmplitudeAnalysis
 - Documentation concerning the theory of operation is available in the Git repository
 - AmpTools "knows" nothing about GlueX data format, physics, etc.
- Issue tracking system on GitHub is used -- report AmpTools problems there if they
 pertain to the core package and not the GlueX implementation



Required Code: Amplitudes

- Provide AmpTools with a method (code via a library) to convert the four-vectors of an event to a complex number
 - inherits from the Amplitude class in AmpTools which defines the interface for the object -- a template is provided to handle some necessary functions
 - see <u>Tutorials/Dalitz/DalitzLib/DalitzAmp/BreitWigner.h</u>
 - accepts arguments as an arbitrarily long list of strings (which will be specified in the config file)
- A collection of GlueX related amplitudes is here:
 - halld_sim/src/libraries/AMPTOOLS_AMPS
- Several optional features:
 - embed floating fit parameters (e.g., BreitWigner mass) into the calculation of the amplitude
 - perform a data reduction step to reduce four-vectors to "user variables," e.g., angles or Lorentz invariants, that are used to compute the amplitude
 - GPU acceleration of amplitude calculation -- requires additional code
 - use of features increases complexity but can optimize performance not a one-size-fits-all solution: ask for advice if you considering additional development to make fits run faster
- The executable you write knows about the existence of the amplitudes through the static registration methods of the AmpToolsInterface
 - AmpToolsInterface::registerAmplitude(BreitWigner());
 - register before creating an instance of AmpToolsInterface to do your fit, generate MC, ...



Required Code: DataReader

- Provide AmpTools with a class that is able to turn a file on disk into a set of four-vectors
 - similar to Amplitude class: inherits from DataReader, uses a template for some key functions, and accepts arguments as a list of strings
 - see example in <u>Tutorials/Dalitz</u>
- Not usually analysis specific, but more specific to the file format
 - GlueX collection of data readers is here: <u>halld_sim/src/libraries/AMPTOOLS_DATAIO</u>
- Common GlueX formats for AmpTools input:
 - standard ROOT tree from tree_to_amptools : ROOTDataReader
 - FSRoot format tree: <u>FSRootDataReader</u>
- Complex functionality can be added:
 - perform filtering or cuts during read into AmpTools
 - bootstrap: random sample with oversampling to evaluate uncertainties
- Not all components of a fit need to use the same data reader
- Like amplitudes, readers need to be registered prior to use:
 - AmpToolsInterface::registerDataReader(DalitzDataReader());



Configuration File: General Remarks

- See sample: <u>Tutorials/Dalitz/run/dalitz3.cfg</u>
- all lines begin with a keyword that informs the parser how to process the rest of the line
 - no continuation character: put it all on one line
 - ordering of the lines is not important
- useful keywords for organizing files:
 - include <file>
 - define <word> (defn1) (defn2) (defn3) ...
 - <word> must be isolated (spaces on each side) to be replaced with one or more words
 - loop <word> <value1> (value2) (value3) ...
 - any line containing <word> will be repeated replacing <word> with <value1>, (value2), ...
 - multiple loops can be in a single line but they must be of the same length N -- then the line is repeated N times stepping through all loops in sync simultaneously
- some special syntax:
 - # as the first character denotes a comment
 - :: is treated like a space
 - [parname] -- use square brackets when the name of a parameter (instead of a numerical value) should be passed as an argument to an amplitude



Reactions, sums, and amplitudes

• Within a reaction, the intensity must be defined as a sum of coherent sums of amplitudes, where each amplitude can be a product of factors

$$egin{aligned} \mathcal{I}(\mathbf{x}) &= \sum_{\sigma} \left| \sum_{lpha} s_{\sigma,lpha} V_{\sigma,lpha} A_{\sigma,lpha}(\mathbf{x})
ight|^2 \ A_{\sigma,lpha}(\mathbf{x}) &= \prod_{\gamma=1}^{n_{\sigma,lpha}} a_{\sigma,lpha,\gamma}(\mathbf{x}), \end{aligned}$$

- ullet Amplitudes can be scaled by a real number $s_{\sigma,lpha}$ (default I) and have a complex production coefficient $V_{\sigma,lpha}$
- This matches the general form for $\eta\pi$ production:

$$I(\Omega, \Phi) = 2\kappa \sum_{k} \left\{ (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m;k}^{(-)} \operatorname{Re}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m;k}^{(+)} \operatorname{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 + P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m;k}^{(+)} \operatorname{Re}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 + P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m;k}^{(-)} \operatorname{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} \right\}.$$

- We absorb $\sqrt{1\pm P_{\gamma}}$ into the definition of Z_l^m and write $[\ell]_m^{(\pm)}$ as either a BreitWigner (for $\ell=2$) or piecewise-defined function (for $\ell=0$): BreitWigner.cc, Zlm.cc, and Piecewise.cc
- Note that one $[\ell]_m^{(\pm)}$ appears in two sums: the production coefficients for each must be constrained to be the same and both terms must be included when computing anything physical from the result
- Multiple reactions are like doing multiple fits simultaneously: contributions to ln(L) add, parameters can be constrained across reactions

Example: Configuring Inputs

fit etapi0_SD_TMD_onePol

define polVal_00 0.3519
define polAngle_00 0.0

define atwo 1.312 0.113

include starting_params.cfg

reaction EtaPi0_00 Beam Proton Eta Pi0

data: signal region events usually with unity weight, may contain backgrounds

bkgnd: (often) weighted sample that is statistically consistent with the background contribution to the signal region (GlueX: weighted RF sidebands + others)

accmc: accepted signal MC, consistent with the signal portion of the data sample (GlueX: remove beam accidentals)

genmc: generated MC, used for denominator in efficiency calculations be mindful of *M*,*t* regions, branching fractions, etc.; (GlueX: this should include the tagger efficiency)

data EtaPi0_00 FSRootDataReader fsroot/tree_pi0eta__B4_M17_M7_DATA_sp17_pol0_SIGNAL_SKIM_A2.root ntFSGlueX_101_1 3

bkgnd EtaPi0_00 FSRootDataReader fsroot/tree_pi0eta__B4_M17_M7_DATA_sp17_pol0_SIDEBANDS_SKIM_A2.root ntFSGlueX_101_1 3 fsroot/tree_pi0eta__B4_M17_M7_DATA_sp17_pol0_SIDEBANDS_SKIM_A2.root.weight ntFSGlueX_101_1_weight weight

accmc EtaPi0_00 FSRootDataReader fsroot/tree_pi0eta__B4_M17_M7_MC_sp17_pol0_SIGNAL_SKIM_A2.root ntFSGlueX_101_1 3 fsroot/tree_pi0eta__B4_M17_M7_MC_sp17_pol0_SIGNAL_SKIM_A2.root.weight ntFSGlueX_101_1_weight weight

genmc EtaPi0_00 FSRootDataReader fsroot/tree_pi0eta__B4_M17_M7_MCGEN_sp17_pol0_GENERAL_SKIM_A2.root ntFSGlueX_101_1 3 MC

reaction: number of particles matches number of four-vectors provided by reader; AmpTools doesn't care about particle names unless they are the same, then the amplitude is symmetrized n.b.: other GlueX software, e.g., gen_amp, may use these names



Example: Setting Up Amplitudes

```
sum EtaPi0 00 ReZ 1-P
                                                                          I(\Omega, \Phi) = 2\kappa \sum_{k} \left\{ \left. (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(-)} \text{Re}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [\ell]_{m; k}^{(+)} \text{Im}[Z_{\ell}^{m}(\Omega, \Phi)] \right|^{2} + (1 - P_{\gamma}) \left| \sum_{\ell, m} [L]_{m; k}^{(+)} \text{Im}[Z
sum EtaPi0 00 ImZ 1+P
sum EtaPi0 00 ReZ 1+P
sum EtaPi0 00 ImZ 1-P
                                                                                                                        (1+P_{\gamma}) \left| \sum_{\ell,m} [\ell]_{m;k}^{(+)} \text{Re}[Z_{\ell}^{m}(\Omega,\Phi)] \right|^{2} + (1+P_{\gamma}) \left| \sum_{\ell,m} [\ell]_{m;k}^{(-)} \text{Im}[Z_{\ell}^{m}(\Omega,\Phi)] \right|^{2} \right\}.
# DEFINE AMPLITUDES
# S-wave amplitudes
amplitude EtaPi0 00::ReZ 1-P::S0- Zlm 0 0 +1 -1 polAngle 00 polVal 00
amplitude EtaPi0_00::ImZ_1+P::S0- Zlm 0 0 -1 +1 polAngle_00 polVal_00
amplitude EtaPi0 00::ReZ 1-P::S0- Piecewise 1.04 1.56 4 23 Neg ReIm [pcwsBin 0ReNeg]
                                                                                                                                                                                                                                                  [pcwsBin 0ImNeg]
 [pcwsBin_1ReNeg] [pcwsBin_1ImNeg] [pcwsBin_2ReNeg] [pcwsBin_2ImNeg] [pcwsBin_3ReNeg]
                                                                                                                                                                                                                                                  [pcwsBin 3ImNeg]
amplitude EtaPi0 00::ImZ 1+P::S0- Piecewise 1.04 1.56 4 23 Neg ReIm [pcwsBin 0ReNeg]
                                                                                                                                                                                                                                                  [pcwsBin 0ImNeg]
 [pcwsBin 1ReNeg] [pcwsBin 1ImNeg] [pcwsBin 2ReNeg] [pcwsBin 2ImNeg] [pcwsBin 3ReNeg]
                                                                                                                                                                                                                                                  [pcwsBin 3ImNeg]
amplitude EtaPi0 00::ImZ 1-P::S0+ Zlm 0 0 -1 -1 polAngle 00 polVal 00
amplitude EtaPi0 00::ReZ 1+P::S0+ Zlm 0 0 +1 +1 polAngle 00 polVal 00
amplitude EtaPi0_00::ImZ_1-P::S0+ Piecewise 1.04 1.56 4 23 Pos ReIm [pcwsBin_0RePos]
                                                                                                                                                                                                                                                  [pcwsBin 0ImPos]
 [pcwsBin_1RePos] [pcwsBin_1ImPos] [pcwsBin_2RePos] [pcwsBin_2ImPos] [pcwsBin_3RePos]
                                                                                                                                                                                                                                                  [pcwsBin 3ImPos]
amplitude EtaPi0_00::ReZ_1+P::S0+ Piecewise 1.04 1.56 4 23 Pos ReIm [pcwsBin_0RePos]
                                                                                                                                                                                                                                                  [pcwsBin 0ImPos]
 [pcwsBin_1RePos] [pcwsBin_1ImPos] [pcwsBin_2RePos] [pcwsBin_2ImPos] [pcwsBin_3RePos]
                                                                                                                                                                                                                                                 [pcwsBin 3ImPos]
# D-wave amplitudes
amplitude EtaPi0 00::ImZ 1-P::a2 D0+ Zlm 2 0 -1 -1 polAngle 00 polVal 00
amplitude EtaPi0 00::ReZ 1+P::a2 D0+ Zlm 2 0 +1 +1 polAngle 00 polVal 00
amplitude EtaPi0 00::ImZ 1-P::a2 D0+ BreitWigner atwo 2 2 3
amplitude EtaPi0 00::ReZ 1+P::a2 D0+ BreitWigner atwo 2 2 3
```

amplitude factors with the same reaction::sum::amplitude are multiplied together



Example: Constraints and Initialization

• We need to constrain the production coefficients $(V_{\sigma,\alpha})$ in front of the specific $[\ell]_m^{(\pm)}$ amplitude that appears in two coherent sums:

```
constrain EtaPi0_00::ImZ_1-P::S0+ EtaPi0_00::ReZ_1+P::S0+
constrain EtaPi0_00::ReZ_1-P::S0- EtaPi0_00::ImZ_1+P::S0-
constrain EtaPi0_00::ImZ_1-P::a2_D0+ EtaPi0_00::ReZ_1+P::a2_D0+
constrain EtaPi0_00::ReZ_1-P::a2_D0- EtaPi0_00::ImZ_1+P::a2_D0-
```

 Initialize (optional) production coefficients with initialized commands -- only need to initialize one of the constrained amplitudes

```
initialize EtaPi0_00::ReZ_1-P::S0- cartesian 1 0 fixed
initialize EtaPi0_00::ReZ_1+P::S0+ cartesian 1 0 fixed
initialize EtaPi0_00::ReZ_1-P::a2_D0- cartesian 27.2 26.9
initialize EtaPi0_00::ReZ_1+P::a2_D0+ cartesian -12.2 19.9
```

- in our example the freedom in the S-wave is in the parameters that make up the piecewise function -- we keep the lead coefficient fixed to one
- Generally required: one of the production coefficients in each coherent sum should be initialized with the flag "real" to avoid overall phase ambiguities
- Parameters are declared and initialized in one line:

```
parameter pcwsBin_0ImNeg -307.092917227354
parameter pcwsBin_0ImPos 423.787950826515
parameter pcwsBin_0ReNeg -29.1570576067053
parameter pcwsBin_0RePos 343.007639956881
parameter pcwsBin_1ReNeg -25.8844360633494
parameter pcwsBin_1RePos 340.647573417975

parameter pcwsBin_1ImPos 0.0 fixed
parameter pcwsBin_1ImNeg 0.0 fixed
```

technical note: these two things together coupled with the fact we are using a Breit-Wigner function in the D wave remove the phase ambiguity



Extending to Multiple Polarization States

- We treat each polarization state as an independent "reaction"
 - data are statistically independent; similar to a coupled-channel or multi-experiment fit, which are ideas supported in AmpTools
 - minimize a global sum of -2 ln(L)
 - constrain amplitudes across reactions up to a floating scale parameter of $\mathcal{O}(1)$ that absorbs differences in integrated luminosity for each data set
 - accepted and generated MC can (likely) be reused for each reaction
- Extend our fit configuration using the AmpTools looping structure:

```
loop LOOPREAC EtaPi0_00 EtaPi0_45 EtaPi0_90 EtaPi0_135
loop LOOPPOLANG polAngle_00 polAngle_45 polAngle_90 polAngle_135
loop LOOPPOLVAL polVal_00 polVal_45 polVal_90 polVal_135
```

which effectively creates four copies of many lines of the config file:

```
reaction LOOPREAC Beam Proton Eta Pi0 amplitude LOOPREAC::ReZ_1-P::S0- Zlm 0 0 +1 -1 LOOPPOLANG LOOPPOLVAL
```

See full example in: etapi0_SD_TMD_allPol.cfg

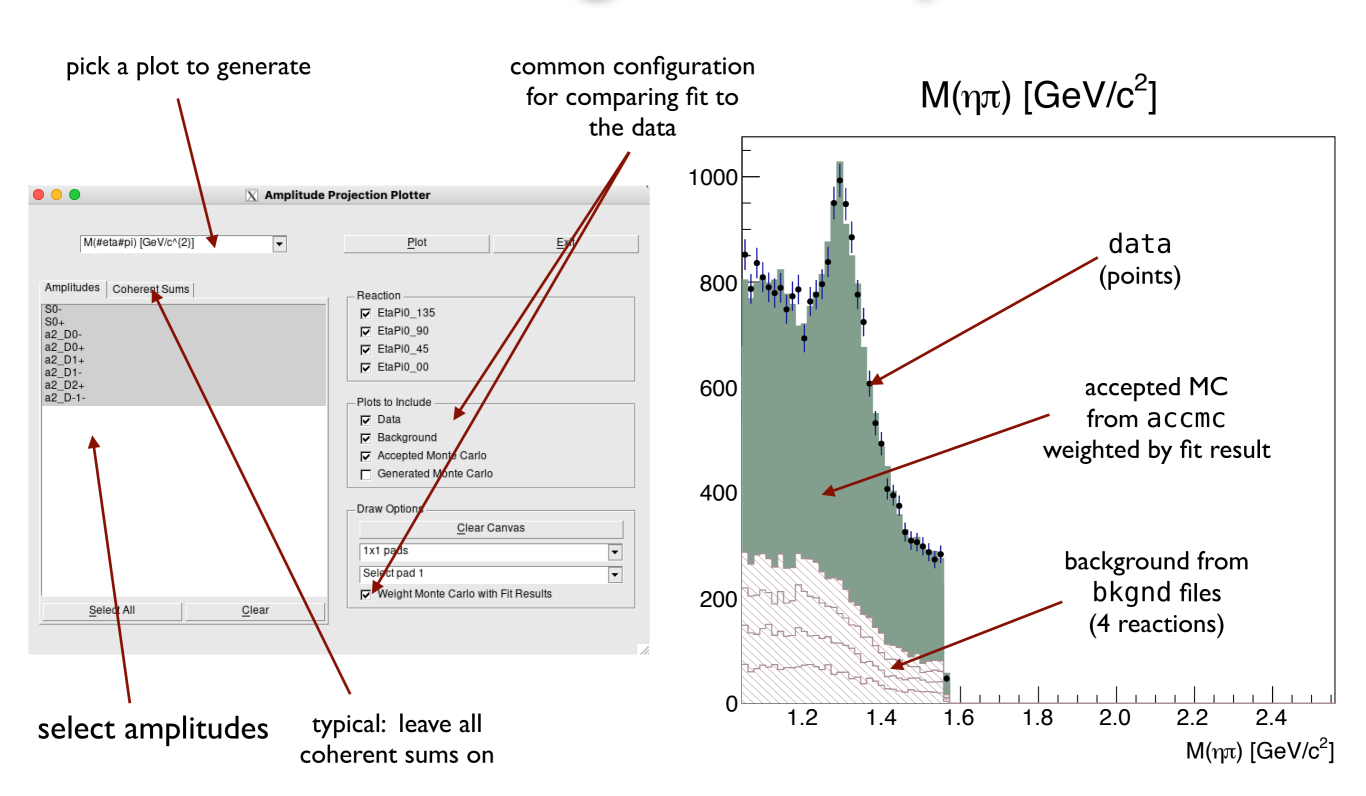


Running the Fit and Visualizing the Result

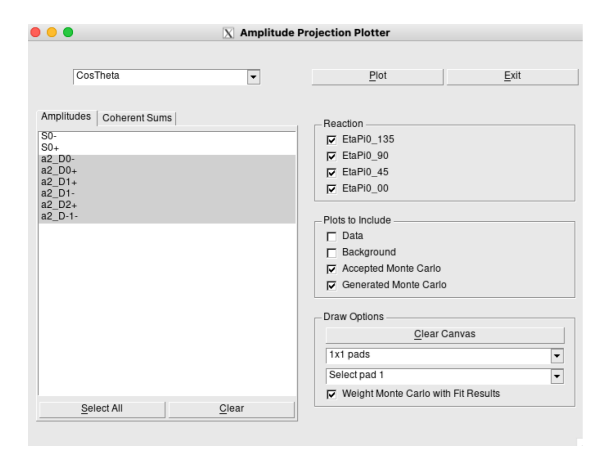
- To run a single core fit interactively invoke fit (source code)
 - run fit -h to see options
 - typical: fit -c etapi0_SD_TMD_allPol.cfg (wait for it... wait for it... ...still waiting...)
 - output: etapi0_SD_TMD_allPol.fit
- Visualization requires the creation of PlotGenerator class to fill histograms
 - see example <u>DalitzPlotGenerator</u> in the AmpTools Dalitz Tutorial
 - collection of GlueX plot generators: <u>halld_sim/libraries/AMPTOOLS_DATAIO</u>
 - for this tutorial see: <u>etaPiPlotGenerator</u>
- And you need an executable that uses this class to generate the plots you want
 - see AmpTools Dalitz Tutorial example: plotResults
 - good for batch processing of many plots -- non-interactive
 - plots of GlueX data that follow were made with: <u>plot_etapi0</u> (in halld_sim)
 - uses AmpPlotter library distributed with AmpTools to provide a GUI front-end to the underlying PlotGenerator
 - good for interactively exploring the amplitudes in a single fit
 - starts ROOT-based XII session: best to run locally.... which, for now, requires a local build of halld sim

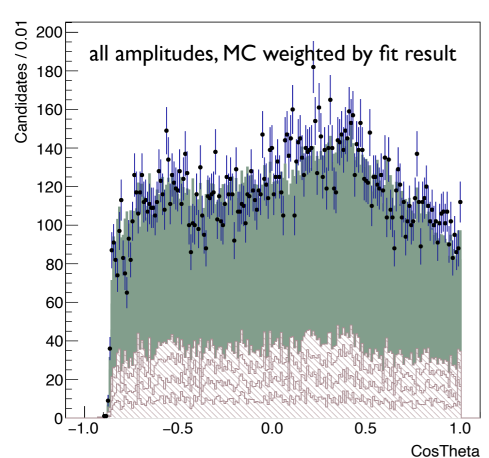


Viewing the Output

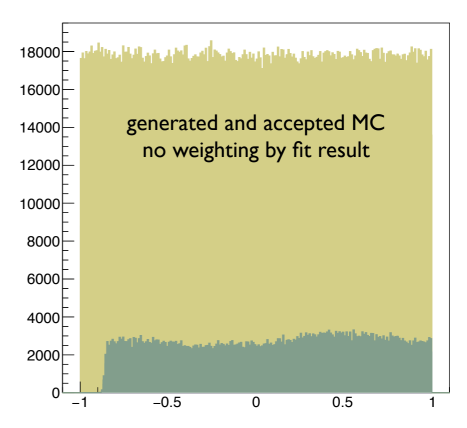




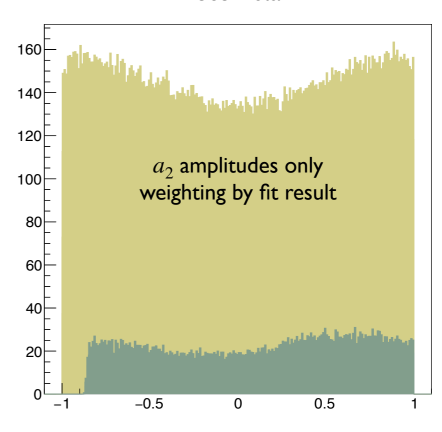




CosTheta



CosTheta



College of Arts and Sciences

Bloomington

Useful Quantities from FitResults

- The *.fit output can be used to create an object of type <u>FitResults</u> which is intended to provide a C++ interface to the results of a fit
- In the session2e directory see <u>displayResults.cc</u> for an example of how to create this object and print some useful information to the screen
 - copy <u>displayResults.cc</u> and the Makefile to your own directory and play with it

```
===== REACTIONS ======
Number of reactions: 4
   (reaction name): (observed signal events from fit) [contribution to −2 ln L]
   EtaPi0 00: 3866.11 +/- 55.0971 [-37262.7]
   EtaPi0 45: 3652.82 +/- 100.789 [-34984.4]
   EtaPi0 90: 4105.35 +/- 109.639 [-38119.9]
   EtaPi0 135: 3748.66 +/- 102.619 [-35453.1]
    TOTAL: 15372.9 +/- 276.389 [-145820]
===== AMPLITUDES =====
   (amplitude name): (acceptance corrected yield) (fit fraction) [detection efficiency]
          : 2350.81 +/- 829.937 (0.0224004) [0.117551]
          : 3781.52 +/- 800.061 (0.0360334) [0.117551]
   a2 D0+
                                                                              Efficiency-corrected number
   a2 D1-
          : 5327.23 +/- 2208.54 (0.0507621) [0.150105]
         : 1198.49 +/- 561.491 (0.0114202) [0.150106]
                                                                         of a_2(1320) events in the data sample
   a2 D-1- : 6256.73 +/- 2144.09 (0.0596192) [0.150105]
                                                                           (assuming \mathcal{B}(a_2 \to \eta \pi^0) = 100 \%
          : 10255.8 +/- 1525.7 (0.0977258) [0.153705]
   50+ : 63909.5 +/- 1078.84 (0.608981) [0.144667]
                                                                               and \mathcal{B}(\eta \to \gamma \gamma) = 100 \%)
          11910.2 +/- 200.964 (0.11349) [0.145053]
    ALL a2(1320) : 29178 +/- 1415.98 (0.278031) [0.147189]
===== MINUIT DIAGNOSTICS ======
  Error Matrix Status ( 3 = full/accurate ): 3
 Last Command Status ( 0 = normal ): 0
  Estimated Distance to Minimum: 5.49816e-06
```

MPI Acceleration

• Invoking fit -c etapi0_SD_TMD_allPol.cfg on an ifarm node will give you time to eat lunch before convergence:

```
MIGRAD evaluation total wall time: 1466.2 s. average time per function call: 199.89 ms.

LIKELIHOOD AFTER MINIMIZATION: -1.4582e+05
```

- AmpTools supports MPI (Message Passing Interface) acceleration: multiple single-threaded processes (potentially across nodes of a cluster) working on a common fit while communicating through the MPI layer
 - allows aggregating RAM across cluster nodes for large data sets
 - enhances speed because each process runs in parallel
 - can be used to accelerate fits on multi-core CPU much like multi-threading a single job
- Requires registration of Amplitudes and parallelized DataReaders (via an AmpTools template) in an executable that can setup and manage the MPI layer:
 - for GlueX use: <u>fitMPI</u> (in halld_sim)
 - if fitMPI is compiled with your release of halld_sim, it is likely you don't need to do anything but load the MPI module in your shell

```
module use /apps/modulefiles
module load mpi/openmpi3-x86_64
```

• full documentation on ifarm/GlueX MPI usage for fitting is here: https://halldweb.jlab.org/wiki/index.php/HOWTO use AmpTools on the JLab farm with MPI



use mpirun to spawn multiple fitMPI processes that work together: mpirun -np 33 fitMPI -c etapi0_SD_TMD_allPol.cfg

top - 10:00:32										
Tasks: 2352 to	tal,	54 1	running,	2268 s	leeping,	27 std	pped	, 2 zomb:	ie	
%Cpu(s): 37.8 ι	us, 3	3.9	sy, 5.5	ni, 52	.5 id, 0	.1 wa,	0.0	hi, 0.0	si, 0.0 st	
KiB Mem : 26380076+total, 641736 free, 49034316 used, 21412470+buff/cache										
KiB Swap: 16777	7212 t	total	1, 15212	576 free	e , 15646 3	36 used	l. 21	381155+ava:	il Mem	
	PR	NI	VIRT	RES	SHR S	%CPU	%MEM	TIME+	COMMAND	
197943 staylor								841:37.07		
210045 shrestha								102:03.46		
129604 shepher		0	906556	200164	101132 R	100.0	0.1	0:34.55	fitMPI	
129610 shepher		0	906552	200152	101128 R	100.0	0.1	0:34.71	fitMPI	
129615 shepher		0	905512	199644	101128 R	100.0	0.1	0:34.71 0:34.65 0:34.62	fitMPI	
129619 shepher		0	905512	199636	101128 R	100.0	0.1	0:34.62	fitMPI	
129620 shepher		0	905512	199640	101128 R	100.0	0.1	0:34.66	fitMPI	
129621 shepher								0:34.62		
129625 shepher								0:34.63		
129633 shepher								0:34.56		
231757 tianye									analysis_R_PVDI	
34907 ebarriga								13:57.64		
92606 ebarriga					113384 R			3:52.32		
109116 ebarriga					113384 R			2:33.11		
129602 shepher					110540 R			0:34.65		
129603 shepher		0			101628 R			0:34.67		
129605 shepher		0			101128 R			0:34.59		
129606 shepher		0			101128 R					
129607 shepher					101124 R					
129609 shepher		0						0:34.67		
129611 shepher		0			101128 R			0:34.64		
129612 shepher		0			101152 R			0:34.58		
129613 shepher		0			101128 R			0:34.64		
129614 shepher		0			101132 R			0:34.64		
129617 shepher		0			101128 R					
129618 shepher					101128 R					
129622 shepher					101128 R					
129623 shepher					101156 R					
129624 shepher					101152 R					
129626 shepher					101128 R			0:34.64		
129627 shepher					101128 R					
129628 shepher					101156 R					
129629 shepher		9			101128 R	99.7	0.1	0:34.58		
129631 shepher		9			101132 R		0.1			
129632 shepher		9			101132 R					
129634 shepher		0			101148 R					
134969 zyy	20	0			13908 R			0:09.12		
205469 jsalvg	20	0			118240 R	99.7			clas12root	
206053 jsalvg	20	9			118224 R	99.7			clas12root	
90143 gluex	20		1606796		58408 S	99.3			•	
129608 shepher		0			101128 R					
129616 shepher		9			101128 R	99.3	0.1			
129630 shepher	d 20	0	905508	199644	101128 R	99.3	0.1	0:34.58	TITMPI	

(single process ifarm CPU)

MIGRAD evaluation total wall time: 1466.2 s. average time per function call: 199.89 ms.

LIKELIHOOD AFTER MINIMIZATION: -1.4582e+05



MIGRAD evaluation total wall time: 65.21 s. average time per function call: 8.8204 ms.

LIKELIHOOD AFTER MINIMIZATION: -1.4582e+05

Do **NOT** do this on the ifarm interactive node as a matter of regular practice!!!

(...but isn't it beautiful to see 33 jobs each at >99% CPU working together on one fit?)

GPU Acceleration

- AmpTools supports acceleration using GPUs (Graphical Processing Units) for numerical computation
 - GPU nodes are available on the ifarm -- these are suitable for fast interactive fitting
- Requires the user to write the amplitude calculation as a GPU "kernel" in a language similar to C called CUDA
 - technically more involved: potential for code-divergence as CPU and GPU code must both be maintained
 - Example in Dalitz Tutorial:
 BreitWigner_kernel.cu
- Many GlueX amplitudes are already implemented in CUDA for GPU fitting
- GPU acceleration is also useful for interactive plotting
- Speed gains are highly dependent on particular fit strategy,
 GPU hardware, etc. -- ask for advice on optimization
- Multiple GPUs can be used (even across nodes) by using the MPI layer where each process uses a GPU

(single process ifarm CPU)

MIGRAD evaluation total wall time: 1466.2 s. average time per function call: 199.89 ms.

LIKELIHOOD AFTER MINIMIZATION: -1.4582e+05



MIGRAD evaluation total wall time: 26.426 s. average time per function call: 3.9056 ms.

LIKELIHOOD AFTER MINIMIZATION: -1.4582e+05

GlueX Documentation: https://halldweb.jlab.org/wiki/index.php/HOWTO use AmpTools on the Lab farm GPUs



Summary

- There is a suite of libraries and executables built off the AmpTools framework for doing amplitude analysis on GlueX data
 - AmpTools is largely independent of physics process or experiment
 - Common GlueX-specific problems to be solved in the context of AmpTools
- A variety of resources are available for issues, questions, help:
 - AmpTools Fits Best Practices
 - AmpTools Issues on GitHub
 - GlueX Software Google group
 - meetings of the Amplitude Analysis Working Group
 - use old technology and phone a friend: mashephe@indiana.edu

