COLLIER a Complex One-Loop LIbrary in Extended Regularizations

Lars Hofer IFAE Barcelona

in collaboration with A. Denner and S. Dittmaier

Los Angeles, June 2015

◆□▶ ◆□▶ ◆□▶ ◆□▶ → □ - のへで

Motivation

- no new particles beyond the SM found so far at the LHC
 new physics might show up only as small deviations from SM predictions
 - \Rightarrow entering precision era of LHC
- perform precise measurements of particle couplings (e.g. couplings of the Higgs boson)
- ► comparison with precise SM predictions ⇒ need SM predictions at NNLO QCD and at NLO electroweak
- one-loop amplitudes needed for NLO virtual and NNLO real-virtual contributions

One-loop amplitudes

general structure of one-loop amplitudes:

$$\int d^{D}q \, \frac{N(q)}{D_{0} \cdots D_{N-1}} = \sum_{r} c_{\mu_{1} \dots \mu_{r}} \underbrace{\int d^{D}q \, \frac{q^{\mu_{1}} \cdots q^{\mu_{r}}}{D_{0} \cdots D_{N-1}}}_{\text{tensor integral } T^{\mu_{1} \dots \mu_{r}}}$$
with $D_{i} = (q + p_{i})^{2} - m_{i}^{2}$

One-loop amplitudes

general structure of one-loop amplitudes:

$$\begin{split} & \underbrace{\int} d^D q \, \frac{N(q)}{D_0 \cdots D_{N-1}} = \sum_r c_{\mu_1 \dots \mu_r} \underbrace{\int} d^D q \, \frac{q^{\mu_1} \cdots q^{\mu_r}}{D_0 \cdots D_{N-1}} \\ & \text{with } D_i = (q+p_i)^2 - m_i^2 \end{split}$$

can be decomposed in terms of scalar integrals:

$$= \sum_{l} d_{l} + \sum_{k} c_{k} + \sum_{j} b_{j} = 0 + \sum_{i} a_{i} + R$$
$$= \sum_{l} d_{l} D_{0}(l) + \sum_{k} c_{k} C_{0}(k) + \sum_{j} b_{j} B_{0}(j) + \sum_{i} a_{i} A_{0}(i) + R$$

▲□▶ ▲□▶ ▲□▶ ▲□▶ ▲□ ● のへぐ

One-loop amplitudes

general structure of one-loop amplitudes:

$$\int d^{D}q \, \frac{N(q)}{D_{0} \cdots D_{N-1}} = \sum_{r} c_{\mu_{1} \dots \mu_{r}} \underbrace{\int d^{D}q \, \frac{q^{\mu_{1}} \cdots q^{\mu_{r}}}{D_{0} \cdots D_{N-1}}}_{\text{tensor integral } T^{\mu_{1} \dots \mu_{r}}}$$

can be decomposed in terms of scalar integrals:

$$= \sum_{l} d_{l} + \sum_{k} c_{k} + \sum_{j} b_{j} = 0 + \sum_{i} a_{i} + R$$
$$= \sum_{l} d_{l} D_{0}(l) + \sum_{k} c_{k} C_{0}(k) + \sum_{j} b_{j} B_{0}(j) + \sum_{i} a_{i} A_{0}(i) + R$$

different approaches for calculation:

w

- ► conventional method (Feynman diagrams) → TI's needed
- generalised unitarity [Ossola, Papadopoulos, Pittau '07, Bern, Dixon, Kosower, Britto, Cachazo, Feng, Ellis, Giele, Melnikov, ...]
- [van Hameren'09; Cascioli, Maierhöfer, Pozzorini'11; Actis, Denner, LH, Scharf, Uccirati'12]

Tools for NLO

- Many tools for NLO calculatios, e.g. FeynCalc/FormCalc, Blackhat, NGluon, aMC@NLO, HELAC-NLO, GoSam, CutTools, HELAC-1LOOP, Samurai, Madloop, OpenLoops, Recola, ...
- Libraries for scalar and tensor integrals, e.g.
 FF [van Oldenborgh], LoopTools [Hahn,Perez-Victoria], QCDLoop [R.K.Ellis,Zanderighi], OneLOop [van Hameren], Golem95C
 [Cullen,Guillet,Heinrich,Kleinschmidt,Pilon,...], PJFry [Fleischer,Riemann]

This talk:

COLLIER = Complex one loop library in extended regularizations

for tran-library for fast and stable numerical evaluation of tensor integrals [Denner,Dittmaier,LH \rightarrow publication in preparation]

Collier: Applications

- successfully used in many calculations of
 - NLO QCD corrections, e.g.
 - $pp \rightarrow t \bar{t} j$ [Dittmaier,Uwer,Weinzierl '07]
 - $pp \rightarrow t \bar{t} b \bar{b}$ [Bredenstein,Denner,Dittmaier,Pozzorini '09]
 - $pp \rightarrow WWb\bar{b}$ [Denner,Dittmaier,Kallweit,Pozzorini '11]
 - $pp \rightarrow WWb\bar{b}H$ [Denner,Feger in prep.] (talk by A.Denner)

NLO EW corrections, e.g.

 $\rm e^+e^- \rightarrow 4~\text{fermions}$ [Denner,Dittmaier,Roth,Wieders '05]

 $pp \rightarrow Hjj$ via VBF [Ciccolini,Denner,Dittmaier '07]

 $pp \rightarrow H + \text{dilepton}$ [Denner,Dittmaier,Kallweit,Mück '11]

 $pp \rightarrow l^+ l^- jj$ [Denner,LH,Scharf,Uccirati '14] (talk by S.Uccirati)

 $pp \rightarrow \mu^+ \mu^- e^+ e^-$ [Biedermann et al. in prep.] (talk by B.Biedermann)

- integrated in automated NLO generators
 - OpenLoops [Cascioli,Maierhöfer,Pozzorini]

(talks by J.Lindert and P.Maierhöfer)

Recola [Actis,Denner,LH,Scharf,Uccirati]

(talk by S.Uccirati)

< ロ > < 団 > < 豆 > < 豆 > < 豆 > < 豆 > < 豆 > < ○ < ○ </p>

Reduction of tensor integrals

Methods implemented in Collier:

applied method depends on number N of propagators

- N = 1, 2: explicit analytical expressions
- N = 3, 4: exploit Lorentz-covariance standard PV reduction [Passarino,Veltman '79]
 + stable expansions in exceptional phase-space regions [Denner,Dittmaier '05]
- ► N ≥ 5: exploit 4-dimensionality of space-time [Melrose '65; Denner,Dittmaier '02,'05; Binoth et al. '05]

Basic scalar integrals from analytic expressions ['t Hooft,Veltman'79; Beenaker,Denner'90; Denner,Nierste,Scharf'91; Ellis,Zanderighi'08; Denner,Dittmaier'11]

< ロ > < 同 > < 三 > < 三 > < 三 > < ○ < ○ </p>

 \Rightarrow fast and stable numerical reduction algorithm

N = 3, 4: PV reduction

►
$$T^{\mu_1...\mu_r} = \int d^D q \, \frac{q^{\mu_1}...q^{\mu_r}}{D_0...D_{N-1}}, \qquad D_i = (q+p_i)^2 - m_i^2$$

contractions:

 $p_i^{\mu}q_{\mu} = -f_i + D_i - D_0, \qquad g^{\mu\nu}q_{\mu}q_{\nu} = m_0^2 + D_0$

 \rightarrow reduction to lower-rank and lower-point integrals

N = 3, 4: PV reduction

►
$$T^{\mu_1...\mu_r} = \int d^D q \, \frac{q^{\mu_1...q^{\mu_r}}}{D_0\cdots D_{N-1}}, \qquad D_i = (q+p_i)^2 - m_i^2$$

contractions:

 $p_i^{\mu}q_{\mu} = -f_i + D_i - D_0, \qquad g^{\mu\nu}q_{\mu}q_{\nu} = m_0^2 + D_0$

 \rightarrow reduction to lower-rank and lower-point integrals

covariant decomposition of tensors:

$$(T^{N})^{\mu_{1}\cdots\mu_{P}} = \sum_{k} \sum_{i_{1},\dots,i_{k}} T^{N,P}_{\underbrace{0\cdots0}_{P-k}} \{\underbrace{g\cdots g}_{(P-k)/2} p_{i_{1}}\cdots p_{i_{k}}\}^{\mu_{1}\cdots\mu_{P}}$$

N = 3, 4: PV reduction

►
$$T^{\mu_1...\mu_r} = \int d^D q \, \frac{q^{\mu_1...q^{\mu_r}}}{D_0\cdots D_{N-1}}, \qquad D_i = (q+p_i)^2 - m_i^2$$

contractions:

 $p_i^{\mu}q_{\mu} = -f_i + D_i - D_0, \qquad g^{\mu\nu}q_{\mu}q_{\nu} = m_0^2 + D_0$

 \rightarrow reduction to lower-rank and lower-point integrals

covariant decomposition of tensors:

$$(T^{N})^{\mu_{1}\cdots\mu_{P}} = \sum_{k} \sum_{i_{1},\dots,i_{k}} T^{N,P}_{\underbrace{0\cdots0}_{P-k}i_{1}\cdots i_{k}} \{\underbrace{g\cdots g}_{(P-k)/2} p_{i_{1}}\cdots p_{i_{k}}\}^{\mu_{1}\cdots\mu_{P}}$$

► system of linear equations for coefficients: → invert for T^{N,P's} ⇒ recursive numerical calculation

$$\Delta T^{N,P} = \left[T^{N,P-1}, T^{N,P-2}, T^{N-1} \right]$$

Gram determinant: $\Delta = \det(Z)$ with $Z_{ij} = 2p_i p_j$

・ロト・西ト・山田・山田・山下

Small Gram determinants

(PV)
$$\Delta T^{N,P} = \left[T^{N,P-1}, T^{N,P-2}, T^{N-1}\right]$$

▲□▶ ▲□▶ ▲□▶ ▲□▶ ▲□ ● のへぐ

small Gram determinant: $\Delta \rightarrow 0$

• $T^{N,P-1}, T^{N,P-2}, T^{N-1}$ become linearly dependent

Small Gram determinants

(PV)
$$\Delta T^{N,P} = [T^{N,P-1}, T^{N,P-2}, T^{N-1}]$$

small Gram determinant: $\Delta \rightarrow 0$

- ► $T^{N,P-1}, T^{N,P-2}, T^{N-1}$ become linearly dependent
- $T^{N,P}$ as sum of $1/\Delta$ -singular terms
 - spurious singularities cancel to give $\mathcal{O}(\Delta)/\Delta\text{-result}$
 - numerical determination of $T^{N,P}$ becomes unstable

▲□▶ ▲□▶ ▲□▶ ▲□▶ ▲□▶ ● のへぐ

Small Gram determinants

(PV)
$$\Delta T^{N,P} = [T^{N,P-1}, T^{N,P-2}, T^{N-1}]$$

small Gram determinant: $\Delta \rightarrow 0$

- ► $T^{N,P-1}, T^{N,P-2}, T^{N-1}$ become linearly dependent
- $T^{N,P}$ as sum of $1/\Delta$ -singular terms
 - ▶ spurious singularities cancel to give $\mathcal{O}(\Delta)/\Delta$ -result
 - numerical determination of $T^{N,P}$ becomes unstable
- ► scalar integrals D₀, C₀, B₀, A₀ become linearly dependent ⇒ O(Δ)/Δ-instabilities intrinsic to all methods relying on the full set of basis integrals D₀, C₀, B₀, A₀
- solution: choose appropriate set of base functions depending on phase-space point

$$\Delta T^{N,P} = \left[T^{N,P-1}, T^{N,P-2}, T^{N-1}\right]$$

◆□ > ◆□ > ◆ □ > ● □ >

$$\Delta T^{N,P+1} = \left[T^{N,P}, T^{N,P-1}, T^{N-1} \right]$$

► exploit linear dependence of T^{N,P}, T^{N,P-1}, T^N for Δ = 0 to determine T^{N,P} up to terms of O(Δ)

< □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □ > < □

$$\Delta T^{N,P+1} = \begin{bmatrix} T^{N,P}, T^{N,P-1}, T^{N-1} \end{bmatrix}$$
$$\Delta T^{N,P+2} = \begin{bmatrix} T^{N,P+1}, T^{N,P}, T^{N-1} \end{bmatrix}$$

► exploit linear dependence of T^{N,P}, T^{N,P-1}, T^N for Δ = 0 to determine T^{N,P} up to terms of O(Δ)

◆□▶ ◆□▶ ◆□▶ ◆□▶ → □ - のへで

• calculate $T^{N,P+1}$ in the same way

$$\Delta T^{N,P+1} = [T^{N,P}, T^{N,P-1}, T^{N-1}]$$

$$\Delta T^{N,P+2} = [T^{N,P+1}, T^{N,P}, T^{N-1}]$$

► exploit linear dependence of T^{N,P}, T^{N,P-1}, T^N for Δ = 0 to determine T^{N,P} up to terms of O(Δ)

< ロ > < 同 > < 三 > < 三 > < 三 > < ○ < ○ </p>

- calculate $T^{N,P+1}$ in the same way
- use $T^{N,P+1}$ to compute $\mathcal{O}(\Delta)$ in $T^{N,P}$

$$\Delta T^{N,P+1} = \left[T^{N,P}, T^{N,P-1}, T^{N-1}\right]$$

$$\Delta T^{N,P+2} = \left[T^{N,P+1}, T^{N,P}, T^{N-1}\right]$$

- ► exploit linear dependence of T^{N,P}, T^{N,P-1}, T^N for Δ = 0 to determine T^{N,P} up to terms of O(Δ)
- calculate $T^{N,P+1}$ in the same way
- use $T^{N,P+1}$ to compute $\mathcal{O}(\Delta)$ in $T^{N,P}$
- ▶ higher orders in ∆ iteratively: O(∆^k)of T^{N,P} requires lower-point T^{N-1} up to rank P + k
- ► basis of scalar integrals effectively reduced (e.g. D₀ from C₀'s)

Coefficients vs. tensors

$$(T^{N})^{\mu_{1}\cdots\mu_{P}} = \sum_{k} \sum_{i_{1},\dots,i_{k}} T^{N,P}_{\underbrace{0\cdots0}_{P-k}} \{\underbrace{g\cdotsg}_{(P-k)/2} p_{i_{1}}\cdots p_{i_{k}}\}^{\mu_{1}\cdots\mu_{P}}$$

of tensor coefficients (TC) vs. # of tensor elements (TE)

	r = 0	r = 1	r = 2	r = 3	r = 4	r = 5	r = 6	
N=3	1	3	7	13	22	34	50	
N = 4	1	4	11	24	46	80	130	#IC < #IE
N = 5	1	5	16	40	86	166	296	
N = 6	1	6	22	62	148	314	610	#TC > #TE
N = 7	1	7	29	91	239	553	1163	1
tensor	1	5	15	35	70	126	210	

Coefficients vs. tensors

$$(T^{N})^{\mu_{1}\cdots\mu_{P}} = \sum_{k} \sum_{i_{1},\dots,i_{k}} T^{N,P}_{\underbrace{0\cdots0}_{P-k}} \{\underbrace{g\cdotsg}_{(P-k)/2} p_{i_{1}}\cdots p_{i_{k}}\}^{\mu_{1}\cdots\mu_{P}}$$

of tensor coefficients (TC) vs. # of tensor elements (TE)

	r = 0	r = 1	r = 2	r = 3	r = 4	r = 5	r = 6	
N=3	1	3	7	13	22	34	50	
N=4	1	4	11	24	46	80	130	#IC < #IE
N = 5	1	5	16	40	86	166	296	
N=6	1	6	22	62	148	314	610	#TC > #TE
N=7	1	7	29	91	239	553	1163	1
tensor	1	5	15	35	70	126	210	

NLO generators OpenLoops and Recola: parametrisation of one-loop amplitude in terms of tensor integrals:

calculated by OpenLoops/Recola

$$\mathcal{M} = \sum_{j} (c^{(j)}_{\mu_1 \dots \mu_{n_j}}) (T^{\mu_1 \dots \mu_{n_j}}_{(j)}) \rightarrow \text{Tensor Integrals}$$

 \Rightarrow need full tensors!

From coefficients to tensors

$$(T^{N})^{\mu_{1}\cdots\mu_{P}} = \sum_{k} \sum_{i_{1},\dots,i_{k}} T^{N,P}_{\underbrace{0\cdots0}_{P-k}} \{\underbrace{g\cdotsg}_{(P-k)/2} p_{i_{1}}\cdots p_{i_{k}}\}^{\mu_{1}\cdots\mu_{P}}$$

In Collier:

- output: coefficients $T^N_{0\cdots 0i_1\cdots i_k}$ or tensors $(T^N)^{\mu_1\cdots \mu_P}$
- efficient algorithm to construct tensors from invariant coefficients for arbitrary N, P via recursive calculation of tensor structures
- for $N \ge 6$: Direct reduction at tensor level



Features of Collier

- complete set of one-loop scalar integrals
- implementation of tensor integrals for (in principle) arbitrary number of external momenta N (tested in many physical processes up to N = 6)
- various expansion methods implemented for exceptional phase-space points (to arbitrary order in expansion parameter)
- mass- and dimensional regularisation supported for IR-singularities
- complex masses supported (unstable particles)
- cache-system to avoid recalculation of identical integrals
- output: coefficients $T^N_{0\cdots 0i_1\cdots i_k}$ or tensors $(T^N)^{\mu_1\cdots \mu_P}$
- ► two independent implementations: COLI+DD

Latest developments

- ▶ validation of 7-point functions in $pp \rightarrow WWb\bar{b}H$ (talk by A.Denner)
- ► improvement in selection of expansion method ⇒ increased stability
- error estimates are performed and returned with the results for the integrals
- improved user friendliness:
 - different in- and output formats supported for calls of tensor integrals
 - various options for output and error handling
 - demo programs illustrating the usage of the library

Structure of Collier



Collier modes

three different modes to run Collier:

- ► 1: Use Coli implementation
- 2: Use DD implementation
- 3: Use Coli and DD and compare results
- ► mode=3:

allows to set parameter check precision:

arguments and results of function calls are reported to an output file if agreement between Coli and DD is worse than check precision

Output of Collier

Structure UV- or IR-singular integrals in $D = 4 - 2\epsilon$ dimensions

$$T^{N} = T_{\rm fin}^{N}(\mu_{\rm UV}^{2}, \mu_{\rm IR}^{2}) + a^{\rm UV}\Delta_{\rm UV} + a_{2}^{\rm IR}\left(\Delta_{\rm IR}^{(2)} + \Delta_{\rm IR}^{(1)}\ln\mu_{\rm IR}^{2}\right) + a_{1}^{\rm IR}\Delta_{\rm IR}^{(1)}$$

・ロト・西ト・モート ヨー うへの

Output of Collier

Structure UV- or IR-singular integrals in $D = 4 - 2\epsilon$ dimensions

$$T^{N} = T_{\rm fin}^{N}(\mu_{\rm UV}^{2}, \mu_{\rm IR}^{2}) + a^{\rm UV}\Delta_{\rm UV} + a_{2}^{\rm IR}\left(\Delta_{\rm IR}^{(2)} + \Delta_{\rm IR}^{(1)}\ln\mu_{\rm IR}^{2}\right) + a_{1}^{\rm IR}\Delta_{\rm IR}^{(1)}$$

► scales μ_{UV}^2 , μ_{IR}^2 and poles $\Delta_{UV} = \frac{c(\epsilon_{UV})}{\epsilon_{UV}}$, $\Delta_{IR,1} = \frac{c(\epsilon_{IR})}{\epsilon_{IR}}$, $\Delta_{IR,2} = \frac{c(\epsilon_{IR})}{\epsilon_{IR}^2}$ can be set to arbitrary real values

 \Rightarrow output of Collier: numerical value for full T^N

Output of Collier

Structure UV- or IR-singular integrals in $D = 4 - 2\epsilon$ dimensions

$$T^{N} = T_{\rm fin}^{N}(\mu_{\rm UV}^{2}, \mu_{\rm IR}^{2}) + a^{\rm UV}\Delta_{\rm UV} + a_{2}^{\rm IR}\left(\Delta_{\rm IR}^{(2)} + \Delta_{\rm IR}^{(1)}\ln\mu_{\rm IR}^{2}\right) + a_{1}^{\rm IR}\Delta_{\rm IR}^{(1)}$$

- ► scales μ_{UV}^2 , μ_{IR}^2 and poles $\Delta_{UV} = \frac{c(\epsilon_{UV})}{\epsilon_{UV}}$, $\Delta_{IR,1} = \frac{c(\epsilon_{IR})}{\epsilon_{IR}}$, $\Delta_{IR,2} = \frac{c(\epsilon_{IR})}{\epsilon_{IR}^2}$ can be set to arbitrary real values \Rightarrow output of Collier: numerical value for full T^N
- ► cancellation of poles can be checked varying Δ_{UV} , $\Delta_{IR,1}$, $\Delta_{IR,2}$
- ► prefactor c(ε) = Γ(1 + ε)(4π)^ε effectively factored out convention can be changed by shifting Δ_{UV}, Δ_{IR,1}, Δ_{IR,2} accordingly
- coefficient a^{UV} of $1/\epsilon_{\text{UV}}$ pole returned also as separate output

Treatment of IR singularities

default: use dimensional regularization

mass regularization supported for collinear singularities:

declare array of squared regulator masses:

 ${\rm minf2}=\{m_1^2,m_2^2,...,m_k^2\}$

with complex (not-necessarily small) numerical values

- if a call of a tensor integral involves an element from minf2, the corresponding mass is
 - set to zero in IR finite integrals
 - kept as regulator mass in IR-singular integrals
- ► In the case of mass regularization the IR-scale µ_{IR} can be interpreted as gluon/photon mass

Error estimates in COLI

Error estimates in Coli: (similar in DD)

1 PV-reduction

error propagation:

 $\delta D_r \sim \max\{ \, \frac{a_r}{\delta D_0}, \frac{b_r}{\delta C_0}, \frac{c_r}{\delta C_{r-1}} \, \}$

with $a_r, b_r \sim 1/\Delta^r, \qquad c_r \sim 1/\Delta$

after calculation: symmetry of coefficients

$$\delta D_r \sim |D_{i_1 i_2 \dots i_r} - D_{i_2 i_1 \dots i_r}|, \quad (0 \neq i_1 \neq i_2 \neq 0)$$

Error estimates in COLI

Error estimates in Coli: (similar in DD)

1 PV-reduction

error propagation:

$$\delta D_r \sim \max\{ a_r \, \delta D_0, b_r \, \delta C_0, c_r \, \delta C_{r-1} \}$$

with $a_r, b_r \sim 1/\Delta^r, \qquad c_r \sim 1/\Delta$

after calculation: symmetry of coefficients

$$\delta D_r \sim |D_{i_1 i_2 \dots i_r} - D_{i_2 i_1 \dots i_r}|, \quad (0 \neq i_1 \neq i_2 \neq 0)$$

2 Expansions: $D_r = D_r^{(0)} + ... + D_r^{(g)}$

• neglected higher orders + error propagation from C's: $\delta D_r = \max\{a_{r,g}, b_r \,\delta C_0, c_g \,\delta C_{r+g}\}$

with $a_{r,g}, c_g \sim \Delta^g$

• extrapolation after calculation: $\delta D_r = D_r^{(g)} \times \frac{D_r^{(g)}}{D^{(g-1)}}$

・ロト・西ト・モン・モー もくの

Precision handling



► target precision: governs selection of expansion method and expansion depth → balancing between precision and run-time

critical precision:

arguments and results of function calls are reported to an output file if estimated accuracy is worse than critical precision

accuracy flag:

stores status of worst integral within all function calls of the same phase space point (reinitialized for new phase-space point)

Choice of reduction scheme in COLI

Strategy for 3-,4-point integrals of rank $r \leq r_{\max}$ in Coli:

(similar in DD)



Cache system

Evaluation of one-loop amplitude leads to multiple calls for the same tensor integral (TI): C(i)

 within one master-call: same TI appears several times in reduction tree



◆□▶ ◆□▶ ▲□▶ ▲□▶ □ のQ@

different master calls and their reductions lead to same TI

Cache system

Evaluation of one-loop amplitude leads to multiple calls for the same tensor integral (TI): C(i)

 within one master-call: same TI appears several times in reduction tree



different master calls and their reductions lead to same TI

Cache system in Collier:

- Identify each TI-call via index pair (N, i):
 - N= number of external master call
 - i = binary index for internal calls (propagated in reduction)
- pointers for each pair (N, i) point to same address in cache if arguments of TI's are identical first call: write cache
 further calls: read cache
- option: cache only internal calls

Conclusions

- Collier= fortran library for numerical calculation of scalar and tensor integrals
- numerical stable results thanks to expansion methods for 3-,4-point integrals
- dimensional and mass regularization supported, as well as complex masses for unstable particles

- two independent implementations: Collier = Coli + DD
- used in NLO generators OpenLoops and Recola
- publication in preparation