

COLLIER
*a Complex One-Loop Library in Extended
Regularizations*

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in collaboration with
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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
⇒ 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:

$$\text{Sun} = \int d^D q \frac{N(q)}{D_0 \cdots D_{N-1}} = \sum_r c_{\mu_1 \dots \mu_r} \underbrace{\int d^D q \frac{q^{\mu_1} \cdots q^{\mu_r}}{D_0 \cdots D_{N-1}}}_{\text{tensor integral } T^{\mu_1 \dots \mu_r}}$$

with $D_i = (q + p_i)^2 - m_i^2$

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with $D_i = (q + p_i)^2 - m_i^2$

can be decomposed in terms of scalar integrals:

$$\begin{aligned} \text{Sun} &= \sum_l d_l \text{Box} + \sum_k c_k \text{Triangle} + \sum_j b_j \text{Bubble} + \sum_i a_i \text{SelfEnergy} + R \\ &= \sum_l d_l D_0(l) + \sum_k c_k C_0(k) + \sum_j b_j B_0(j) + \sum_i a_i A_0(i) + R \end{aligned}$$

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different approaches for calculation:

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

Tools for NLO

- ▶ Many tools for NLO calculations, 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

fortran-library for fast and stable numerical evaluation of
tensor integrals [Denner,Dittmaier,LH → 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.
 - $e^+e^- \rightarrow 4$ 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)

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]

▶ $N \geq 5$: 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]

⇒ **fast and stable numerical reduction algorithm**

$N = 3, 4$: PV reduction

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

contractions:

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

→ reduction to **lower-rank** and **lower-point** integrals

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→ reduction to **lower-rank** and **lower-point** integrals

► covariant decomposition of tensors:

$$(T^N)^{\mu_1 \dots \mu_P} = \sum_k \sum_{i_1, \dots, i_k} \underbrace{T_{0 \dots 0}_{i_1 \dots i_k}^{N, P}}_{P-k} \left\{ \underbrace{g \dots g}_{(P-k)/2} p_{i_1} \dots p_{i_k} \right\}^{\mu_1 \dots \mu_P}$$

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► system of linear equations for coefficients:

→ invert for $T^{N, P}$'s ⇒ recursive numerical calculation

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

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

Small Gram determinants

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

small Gram determinant: $\Delta \rightarrow 0$

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

Small Gram determinants

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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**

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 - ▶ spurious singularities cancel to give $\mathcal{O}(\Delta)/\Delta$ -result
 - ▶ numerical determination of $T^{N,P}$ becomes **unstable**
- ▶ scalar integrals D_0, C_0, B_0, A_0 become linearly dependent
 $\Rightarrow \mathcal{O}(\Delta)/\Delta$ -instabilities **intrinsic** to all methods relying on the full set of basis integrals D_0, C_0, B_0, A_0
- ▶ **solution**: choose **appropriate set** of base functions depending on phase-space point

Expansion in Gram determinant

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

Expansion in Gram determinant

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

- ▶ exploit linear dependence of $T^{N,P}, T^{N,P-1}, T^N$ for $\Delta = 0$ to determine $T^{N,P}$ up to terms of $\mathcal{O}(\Delta)$

Expansion in Gram determinant

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



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

- ▶ exploit linear dependence of $T^{N,P}, T^{N,P-1}, T^N$ for $\Delta = 0$ to determine $T^{N,P}$ up to terms of $\mathcal{O}(\Delta)$
- ▶ calculate $T^{N,P+1}$ in the same way

Expansion in Gram determinant

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



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

Expansion in Gram determinant

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

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

- ▶ exploit linear dependence of $T^{N,P}, T^{N,P-1}, T^N$ for $\Delta = 0$ to determine $T^{N,P}$ up to terms of $\mathcal{O}(\Delta)$
- ▶ 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 Δ iteratively:
 $\mathcal{O}(\Delta^k)$ of $T^{N,P}$ requires lower-point T^{N-1} up to rank $P+k$
- ▶ basis of **scalar integrals** effectively reduced
(e.g. D_0 from C_0 's)

Coefficients vs. tensors

$$(T^N)^{\mu_1 \dots \mu_P} = \sum_k \sum_{i_1, \dots, i_k} T_{\underbrace{0 \dots 0}_{P-k} i_1 \dots i_k}^{N, P} \underbrace{\{g \dots g\}}_{(P-k)/2} p_{i_1} \dots p_{i_k} \}^{\mu_1 \dots \mu_P}$$

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	#TC < #TE
$N = 4$	1	4	11	24	46	80	130	
$N = 5$	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	
tensor	1	5	15	35	70	126	210	

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NLO generators **OpenLoops** and **Recola**:

parametrisation of one-loop amplitude in terms of **tensor integrals**:

calculated by OpenLoops/Recola

$$\mathcal{M} = \sum_j c_{\mu_1 \dots \mu_{n_j}}^{(j)} T_{(j)}^{\mu_1 \dots \mu_{n_j}} \rightarrow \text{Tensor Integrals}$$

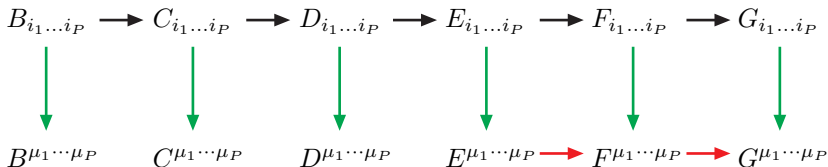
⇒ need full tensors!

From coefficients to tensors

$$(T^N)^{\mu_1 \dots \mu_P} = \sum_k \sum_{i_1, \dots, i_k} \underbrace{T_{0 \dots 0 i_1 \dots i_k}^{N, P}}_{P-k} \underbrace{\{g \dots g p_{i_1} \dots p_{i_k}\}}_{(P-k)/2}^{\mu_1 \dots \mu_P}$$

In Collier:

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



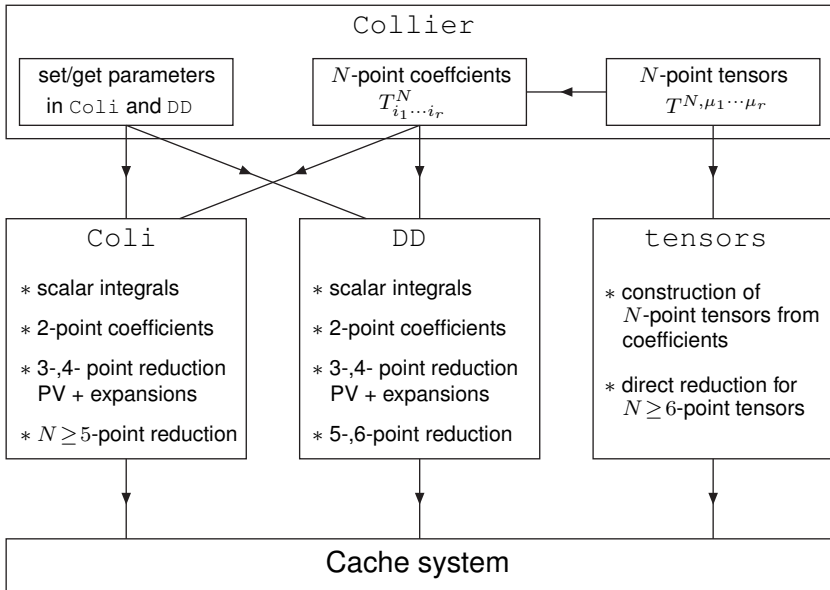
Features of Collier

- ▶ 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 $T_{0\dots 0i_1\dots i_k}^N$ or tensors $(T^N)^{\mu_1\dots\mu_P}$
- ▶ 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

$$T^N = T_{\text{fin}}^N(\mu_{\text{UV}}^2, \mu_{\text{IR}}^2) + a^{\text{UV}} \Delta_{\text{UV}} + a_2^{\text{IR}} \left(\Delta_{\text{IR}}^{(2)} + \Delta_{\text{IR}}^{(1)} \ln \mu_{\text{IR}}^2 \right) + a_1^{\text{IR}} \Delta_{\text{IR}}^{(1)}$$

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► scales

$$\mu_{\text{UV}}^2, \quad \mu_{\text{IR}}^2$$

and poles $\Delta_{\text{UV}} = \frac{c(\epsilon_{\text{UV}})}{\epsilon_{\text{UV}}}$, $\Delta_{\text{IR},1} = \frac{c(\epsilon_{\text{IR}})}{\epsilon_{\text{IR}}}$, $\Delta_{\text{IR},2} = \frac{c(\epsilon_{\text{IR}})}{\epsilon_{\text{IR}}^2}$

can be set to arbitrary real values

⇒ output of Collier: numerical value for full T^N

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⇒ output of Collier: numerical value for full T^N

- ▶ cancellation of poles can be checked varying Δ_{UV} , $\Delta_{\text{IR},1}$, $\Delta_{\text{IR},2}$
- ▶ prefactor $c(\epsilon) = \Gamma(1 + \epsilon)(4\pi)^\epsilon$ effectively factored out
convention can be changed by shifting Δ_{UV} , $\Delta_{\text{IR},1}$, $\Delta_{\text{IR},2}$
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:

$$\text{minf2} = \{m_1^2, m_2^2, \dots, m_k^2\}$$

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:

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

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

- ▶ after calculation: **symmetry of coefficients**

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

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

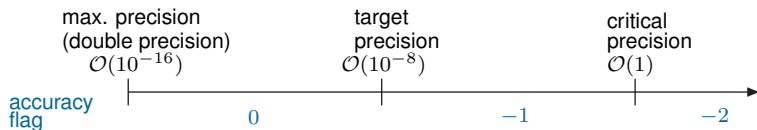
- ▶ **neglected higher orders** + **error propagation from C's**:

$$\delta D_r = \max\{a_{r,g}, b_r \delta C_0, c_g \delta C_{r+g}\}$$

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

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

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)

1 PV reduction:

accuracy for rank r_{\max}
better than target precision?

yes
→

use PV reduction
for $r \leq r_{\max}$ done

↓ no

2 Expansions:

do $g = 0, g_{\max}$
accuracy for rank r_{\max}
and expansion up to order g
better than target precision?
end do

yes
→

use expansion
up to order g done
for $r \leq r_{\max}$

↓ no

3 No method optimal:

do $r_0 = r_{\max}, 0$
is there a method with
better accuracy for rank r_0 ?
end do
done

→

use for $r \leq r_{\max}$ method
with best accuracy for r_{\max}

↙

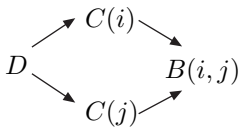
↔

use this method
for $r \leq r_0$

Cache system

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

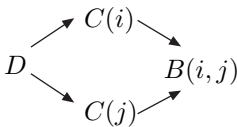
- ▶ **within one master-call:**
same TI appears several times
in reduction tree
- ▶ **different master calls** and their reductions lead to same TI



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- ▶ **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**