# Semi-Automatic Floating-Point Implementation of Special Functions

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*Abstract*—This work introduces an approach to the computerassisted implementation of mathematical functions geared toward special functions such as those occurring in mathematical physics. The general idea is to start with an exact symbolic representation of a function and automate as much as possible of the process of implementing it.

In order to deal with a large class of special functions, our symbolic representation is an implicit one: the input is a linear differential equation with polynomial coefficients along with initial values. The output is a C program to evaluate the solution of the equation using domain splitting, argument reduction and polynomial approximations in double-precision arithmetic, in the usual style of mathematical libraries.

Our generation method combines symbolic-numeric manipulations of linear ODEs with interval-based tools for the floatingpoint implementation of "black-box" functions. We describe a prototype code generator that can automatically produce implementations on moderately large intervals. Implementations on the whole real line are possible in some cases but require manual tool setup and code integration. Due to this limitation and as some heuristics remain, we refer to our method as "semiautomatic" at this stage.

Along with other examples, we present an implementation of the Voigt profile with fixed parameters that may be of independent interest.

## I. INTRODUCTION

Fixed-precision floating-point ("FP") operations come with varying performance and accuracy guarantees. Basic operations such as multiplication are typically implemented in hardware and, being correctly rounded, provide perfect accuracy. So-called elementary functions like exp and log are provided in software through general-purpose mathematical libraries (libms). They are well-optimized for performance and either correctly rounded, as recommended by the IEEE754 Standard [16], or provided with maximum error not exceeding one unit in the last place.

This work is concerned with special functions, i.e., "functions which are widely used in scientific and technical applications, and of which many useful properties are known" [12]. Some, like Bessel functions or the Gaussian error function, are present in some libms. Most are provided by specialized libraries such as GSL or Root. Others are only available in computer algebra systems like Maple and Pari and have no efficient fixed-precision implementation. The NIST *Digital Library of Mathematical Functions* [1] contains a good overview of existing implementations.

There is a huge gap in both performance and rigor between these implementations and state-of-the-art elementary functions. However, implementing or reimplementing special functions manually in the classical way is tedious and requires considerable expertise. And indeed, in many cases, publicly available implementations of a given special function all derive from a single source, typically Cephes [23].

This article presents the first results of a project exploring the implementation of special functions through automatic *code generation*. In our view, this is the only feasible way to implement wider classes of special functions while bridging the quality gap with respect to elementary functions. Additionally, code generation allows for production of specialized implementations taylored to specific applications, in terms of accuracy, supported domain or code properties.

Our focus is on special functions satisfying linear ordinary differential equations of the form

$$p_r(x)f^{(r)}(x) + \dots + p_1(x)f'(x) + p_0(x)f(x) = 0, \quad p_i \in \mathbb{Q}[x].$$
 (1)

Such functions are called *D*-finite or holonomic. A key idea in the field of D-finite functions is that an equation such as (1) along with initial values makes up a concise exact representation of its solution that can be used in computations.

Many common special functions can be defined this way. Among the better known special functions of mathematical physics, univariate D-finite functions include in particular the error function (and related functions such as the Dawson functions or the Voigt profile), the Airy functions (as well as Scorer functions, generalized Airy functions...), the Bessel and Hankel functions, the Struve functions of integer order, the hypergeometric and generalized hypergeometric functions, the spheroidal wave functions, and the Heun functions.

Under the term *code generator*, we understand a software tool which takes as input a description of a function f, a set of floating-point inputs X and a target accuracy  $\varepsilon > 0$ , and which generates source code (in our case, in C) that provides a function  $\tilde{f}$  satisfying

$$\forall x \in X, \quad \left| \frac{\tilde{f}(x) - f(x)}{f(x)} \right| \le \varepsilon.$$
 (2)

In this work, we consider inputs in IEEE754 double precision (binary64) format taken in a relatively small interval [a, b], say of width b - a < 100, and accuracies compatible with that format, i.e.,  $2^{-53} \le \varepsilon \le 1$ . These restrictions can be loosened

with some manual work so that implementations on the whole set of double precision numbers become feasible.

Code generation for mathematical functions is not entirely a new idea. A powerful toolbox, including in particular Gappa [8] and Sollya [6], has been developed in recent years to help human developers with the most tedious steps of the implementation process. Building on these advances, several authors have considered code generation both for "flavors" of a fixed function and for "black-box" functions given as executable code [19], [9], [3], [18], [4]. Independently, Beebe [2] produced a new, very extensive mathematical library with the help of semi-automatic code generation. We base our work on the black-box approach of [19], where the starting point is executable code able to evaluate f and its first few derivatives on intervals, up to any required precision. Providing such an evaluator is easy when f is a composition of basic elementary functions, thanks to multiple-precision interval libraries, but it is a major limitation for more general functions.

Nevertheless, if f satisfies some kind of functional equation, any rigorous arbitrary-precision numerical solver could in principle play the rôle of the black box. In the case of D-finite functions, such solvers can be built in practice [25], [31]. Moreover, the behavior of D-finite functions in the neighborhood of their singularities is well-understood, opening the door to further extensions of the method. These observations were, in some sense, the starting point of the present work. We soon observed that the interval black-box model it is not well suited to our case, as the number of queries to the function tends to be very large. This led us to favor a different interface, based on polynomial approximations, as described in Section II.

This article is organized as follows. Section II describes the general architecture of our function generation pipeline and introduces our prototype implementation. Section III focuses on the frontend part, that is, the rigorous ODE solver. Section IV discusses the backend, which is the part in charge of floating-point implementation choices. The remaining section presents experimental results and applications.

## II. OVERALL ARCHITECTURE

This text describes both an approach to code generation for D-finite functions and experiments performed with a prototype implementation of this approach. Our prototype, named Frankenstein, is available from https://gforge.inria.fr/ git/metalibm/frankenstein.git. It is based on modified versions of two existing software packages, Metalibm [18], written in Sollya [6], and NumGfun [21], written in Maple. The name should give a pretty accurate idea of how the combination is realized. Thus, not all design choices have the same significance: while some would survive a cleaner rewrite, others are only justified by the goal of producing a usable prototype out of two research-quality codes that had never been intended to work with each other. Here we focus on the former category, with brief mentions of more peculiar features of Frankenstein as necessary.

Consider a sufficiently small interval  $I = [a, b] \subset \mathbb{R}$  and a solution f of (1) defined on I. Assume for simplicity that

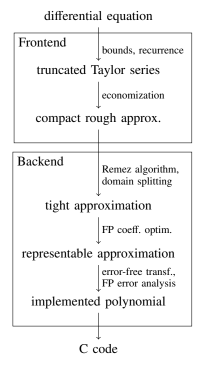


Figure 1. A piecewise polynomial approximation pipeline.

 $0 \in I$  and  $p_r(x) \neq 0$  for  $x \in I$ . The function f is then analytic on I, and the vector of initial values  $F(0) = (f(0), \ldots, f^{(r-1)}(0))$ characterizes f among the solutions of (1). Assume additionally that  $|f(x)| \leq 2^{1024}(1 - 2^{-53})$  for  $x \in I$ . Then, special floatingpoint values (NaN,  $\pm \infty$ ) occur neither on input nor on output. Denoting by  $\mathbb{F}$  the set of double-precision numbers, our goal is to produce a program computing a value  $\tilde{f}(x) \in \mathbb{F}$  satisfying (2) with  $X = I \cap \mathbb{F}$ .

Our method takes as input  $(p_i)_{i=0}^r$  and F(0) (which together define f), I,  $\varepsilon$ , and various constraints on allowable implementations. It either succeeds and produces a program satisfying (2) by construction<sup>1</sup>, or fails if no implementation fitting the constraints is found. Failure does not imply that no feasible solution exists.

One way to view the process leading from the definition of f by (1) to an implementation is as a rigorous piecewise polynomial approximation pipeline, as illustrated on Figure 1. Each stage receives a set of approximations of f (and possibly its first few derivatives) by polynomials on subintervals of Iand produces approximations with different properties that are passed on to subsequent stages. The subintervals vary from stage to stage, and typically overlap even within a single stage. For example, the domain splitting procedure can use both a rough approximation on the whole domain to get a general picture of the behavior of f, and tight approximations on tiny intervals to compute precise values.

The complete pipeline can be approximately divided into a frontend and a backend. Roughly speaking, the frontend is the

<sup>&</sup>lt;sup>1</sup>This is not entirely true of Frankenstein yet. Although the overwhelming part of the code uses rigorous error bounds, some heuristic estimates remain.

part where decisions are driven by the analytic properties of f. The backend is the part where they are driven by the floatingpoint environment and other implementation constraints. To work with another class of mathematical functions, one would replace the frontend; to target a different evaluation environment (fixed-point arithmetic, say), one would swap out the backend. There is no formal abstraction of how the backend can query the function, though, and both parts have full knowledge of the implementation problem.

In Frankenstein, the frontend and the backend respectively correpond to NumGfun and Metalibm. For convenience reasons, the process is driven by the backend, which can ask the frontend for approximations of variable quality on various subintervals. As we reuse code written for the black-box function model, the backend mostly uses these approximations to perform interval evaluations, but we expect to make more direct use of the polynomial representation in future developments.

Our general implementation strategy fits the standard pattern of special function implementations and is especially close to that of Harrison [14]. Classical argument reduction algorithms based on algebraic properties of elementary functions typically do not apply. Accordingly, the only feature of f that we consider to reduce the implementation domain is its parity. Parity properties can be detected either by numerical comparison of f(x) and f(-x) (as Frankenstein currently does), or based on the differential equation. Periodic functions could be handled in a similar way but are very uncommon.

We then single out a small number of "interesting points", the neighborhood of which need to be handled in a special way. Under our working hypotheses, interesting points include x = 0, where the floating-point grid is denser than usual, and the points x with  $f(x) \approx 0$ , where obtaining a relative error bound requires special care. In a more general setting, one would add at least  $\pm \infty$  and the singularities of f to the list. In a small interval around each interesting point, approximations of f are computed and implemented in a way that takes into account the special constraints. The remaining subintervals (if any) are further subdivided until approximation by small-degree polynomials becomes feasible.

Let us now consider in more detail the main ingredients of the process, starting with the frontend.

#### III. FROM A DIFFERENTIAL EQUATION TO RIGOROUS APPROXIMATIONS

The frontend's rôle is to provide the backend with rigorous polynomial approximations of f of the form

$$\forall x \in J, \quad |f(x) - p(x)| \le \eta \tag{3}$$

for various subintervals  $J \subseteq I$ . As Equation (3) suggests, in our architecture, the approximations come with *absolute* error bounds. Indeed, these are often easier to obtain and this choice does not hinder the use of relative error bounds in later stages.

Values of J above can range from point intervals  $J = \{x\}$  to J = I. It is not necessary that the frontend be able to find approximations on arbitrarily wide intervals, as the backend has the necessary logic to split J if the returned error bound is infinite (or too large). The only requirement is that sufficiently

precise queries on sufficiently thin intervals eventually produce satisfying approximations.

The smallest useful value of  $\eta$  in our setting is slightly less than the smallest subnormal number,  $2^{-1074}$ . In principle, it would be possible to replace f once and for all by a set of polynomial approximations achieving such an accuracy. The frontend of Frankenstein is more flexible and can compute arbitrarily good approximations. Precise polynomial approximations on wide intervals quickly become large and costly to compute. For historical reasons, our frontend provides a separate interface dedicated to points and very thin intervals that always returns "polynomial approximations" reduced to constants, which can reach absolute precisions  $\log \eta^{-1}$  in the hundred of thousands if necessary.

The computation of the approximations (3) is based on a combination of classical techniques that we now summarize. We refer the reader to [31], [21] for details.

## A. Transition matrices

Recall that f is specified by the differential equation (1) and the vector of initial values F(0), or, to be precise, rigorous arbitrary precision approximations of F(0). Let us also extend the notation  $F(x) = (f(x), \ldots, f^{(r-1)}(x))$  to arbitrary x. Polynomial approximations on an interval  $J \subseteq I$  will be derived from the Taylor expansion of f around the center c of J. This expansion is easily computed from the differential equation and the "initial value" F(c). To deduce F(c) from F(0), we use Taylor expansions at intermediate points  $x_0 = 0, x_1, \ldots, x_n = c$  chosen in such a way that  $x_{k+1}$  lies within the disk of convergence of the Taylor expansion of F at  $x_k$ . In other words, our frontend is essentially a rigorous ODE solver based on the so-called method of Taylor series [20].

More precisely, we proceed as follows [7], [31]. By linearity of (1), for all  $x, y \in I$ , there exists a matrix  $\Delta_x(y) \in \mathbb{R}^{r \times r}$ depending only on (1) (not on the particular solution f) and such that  $F(y) = \Delta_x(y) F(x)$ . We have  $\Delta_x(z) = \Delta_y(z) \Delta_x(y)$  for all x, y, z, and hence

$$F(c) = \Delta_0(c)F(0) = \Delta_{x_{n-1}}(c) \dots \Delta_{x_1}(x_2) \Delta_0(x_1) F(0).$$
(4)

The entries of  $\Delta_x(y)$  are values at *y* of solutions of (1) corresponding to unit initial values at *x* (or derivatives thereof). Denoting by  $\rho(x)$  the distance from *x* to the nearest complex root of the leading coefficient  $p_r$  of (1), the Taylor expansion at *x* of any solution of (1) has radius of convergence at least  $\rho(x)$ . It is a classical fact that the coefficients of these expansions obey linear recurrences with polynomial coefficients, making it easy to compute as many terms as needed. As we assumed that  $p_r$  does not vanish on *I*, computing *F*(*c*) reduces to forming the product (4), where each factor  $\Delta_x(y)$  can be evaluated by summing convergent power series whose coefficients are easy to compute. Binary splitting can be used to compute the partial sums efficiently to very high precisions [7].

Our assumption that the initial values are provided at a point of *I* is artificial: the above argument still works with  $0 \notin I$ as long as the leading coefficient  $p_r(x)$  of (1) does not vanish between 0 and *I*. In addition, (1) actually defines *f* for complex values of *x*, and nothing prevents the path  $(x_0, \ldots, x_n)$  from going through the complex plane. If  $p_r(s) = 0$  for some *s* between 0 and *I*, analytic continuation along a path avoiding *s* still defines *f* on *I* (in general in a way that depends on the path). Furthermore, when  $0 \notin I$ , we can relax the assumption that  $p_r(0) \neq 0$ : the theory extends to the case where 0 is a so-called regular singular point of (1) [32], [21]. Both of these extensions are supported in Frankenstein, albeit with specific tool setup and subject to heuristic error estimates in some cases. The second one is useful because many classical special functions are best characterized by their behavior at regular singular points of their defining equation. A typical example is the family of Bessel functions (see Example 3 below).

## B. Error bounds

It is crucial that the frontend provides rigorous error bounds on the approximations it computes, as these approximations are the backend's only access to the function f. Here, we recall a simple bound computation technique based on the theory of majorizing series [15, Chap. 2]. Frankenstein (via NumGfun) actually uses a more sophisticated variant of the same technique [22], but the present version conveys the main ideas and would probably suffice for our purposes.

Consider a power series with matrix coefficients (or, equivalently, a matrix of power series)  $Y(x) = \sum_{n=0}^{\infty} Y_n x^n \in \mathbb{R}^{r \times r}[[x]]$ , and let  $\|\cdot\|$  denote a matrix norm. We write  $Y \leq w$  if  $w(x) = \sum_{n=0}^{\infty} w_n x^n \in \mathbb{R}_+[[x]]$  is a power series that bounds *Y* coefficient-wise, i.e.,  $\|Y_n\| \leq w_n$  for all  $n \in \mathbb{N}$ . The method transfers such bounds on the coefficients of differential equations to similar bounds on the solutions.

Our goal is to control the error committed by truncating the Taylor expansions of a solution of (1). Without loss of generality, we restrict ourselves to Taylor expansions at the origin. Rewrite (1) in matrix form, as

$$Y'(x) = P(x)Y(x),$$
(5)

where  $P \in \mathbb{Q}(x)^{r \times r}$  is a companion matrix with entries of the form  $p(x)/p_r(x)$ , and for notational simplicity *Y* is also taken to be a matrix. The matrix function *P* admits a convergent power series expansion at 0, whose coefficients  $P_n$  satisfy  $||P_n|| = O(\alpha^n)$  for all  $\alpha > \rho(0)^{-1}$ . Given such an  $\alpha$ , it is not too hard to compute M > 0 such that

$$P(x) \leq q(x) := \frac{\alpha M}{(1 - \alpha x)}.$$

Expanding both sides of (5) in power series and collecting the matching powers of x, we obtain

$$(n+1)Y_n = \sum_{j=0}^n P_n Y_{n-j}.$$
 (6)

The same reasoning applied to the equation w'(x) = q(x)w(x) yields

$$(n+1)w_n = \sum_{j=0}^n q_n w_{n-j}.$$
 (7)

Now assume  $w_0 \ge ||Y_0||$ . Comparing (6) with (7), we see by induction that  $Y(x) \le w(x)$ . But the second equation is solvable in closed form: we have

$$w(x) = \frac{w_0}{(1 - \alpha x)^M}.$$

This explicit expression makes it easy to bound the tails  $w_n x^n + w_{n+1}x^{n+1} + \cdots$ , and hence also  $Y_n x^n + Y_{n+1}x^{n+1} + \cdots$ , which yields truncation orders that guarantee a certain accuracy.

# C. Polynomial approximations

We are now ready to combine the results of the previous sections to obtain rigorous polynomial approximations on a given interval J. Write  $J = [c - \delta, c + \delta]$ , and assume that  $\delta < \rho(c)$ . With the notation of Section III-A, we have

$$F(c+\xi) = \Delta_c(c+\xi) \,\Delta_{x_{n-1}}(c) \dots \Delta_{x_1}(x_2) \,\Delta_0(x_1) \,F(0) \tag{8}$$

for  $|\xi| \le \delta$ . Each factor (except F(0)) is given by power series that we can truncate so as to guarantee a prescribed accuracy. Error bounds on the individual factors are combined by repeated use of the inequality

$$\|\tilde{A}\tilde{B} - AB\| \le \|\tilde{A}\| \|\tilde{B} - B\| + \|\tilde{A} - A\| \|B\|$$

Once we replace each series by a truncation (and F(0) by an approximation with rational entries), the entries of (8) become polynomials in  $\xi$  with rational coefficients.

Thus, we compute the entries of  $\Delta_c(c + \xi)$  truncated to a suitable order as polynomials in  $\xi$  and multiply the resulting matrix by an approximation of  $\Delta_0(c) F(0)$ . The entries of the result readily provide the first *r* derivatives of *f*. Higher-order derivatives, if needed, can be obtained by multiplying  $F(c + \xi)$  on the left by a row vector of rational functions (or polynomial approximations of rational functions) deduced from (1). In Frankenstein, most of these steps are performed exactly, using multiple precision rational arithmetic. Roundoff errors in the remaining steps are taken into account in the result.

Taylor series typically do not provide good approximations on intervals. Additionally, the bounds of Section III-B can be quite pessimistic. For these reasons, polynomials computed as outlined above tend to have very high degree. However, due to the way they are constructed, the rescaled coefficients  $c_n \delta^n$ quickly decrease to zero, and typically  $|c_n \xi^n| \ll \eta$  for large *n*. Before handing it to the backend, the frontend hence reduces the degree of the computed polynomial *p* by *economization* [11, §4.6]: while the leading term of p is small compared to  $\eta$ , a multiple of the Chebyshev polynomial  $T_n$  (rescaled to the interval J) chosen so that the leading terms cancel is subtracted from p. The error bound is updated accordingly. (The choice of Chebyshev polynomials makes it possible to take advantage of the fact that the error bound only needs to hold for  $x \in J$ , not for complex x with  $|x - c| < \delta$ .) This procedure is very effective at producing polynomials of reasonable size that can easily be manipulated by the backend.

## IV. FROM RIGOROUS APPROXIMATIONS TO EVALUATION CODE

It is the backend's job to transform the high-degree, rough approximation produced by the frontend into fine-tuned approximations with suitable FP properties and eventually, into source code. In absence of classical argument reduction techniques, implementation of the function is reduced to piecewise polynomial approximation. The backend hence needs to compute four pieces of information, as discussed in the next subsections.

1) The implementation interval I needs to be split into subintervals  $I_k$ , together covering I, such that an approximation polynomial of small degree is possible over each  $I_k$ .

2) These small degree polynomials need to be computed, initially with real coefficients.

3) Connected with the approximation step is the problem of choosing an appropriate translation  $f(t_k + \cdot)$  on a new domain  $I_k - t_k$ . Indeed, the evaluation of the approximation behaves better when  $I_k - t_k$  is a small interval around 0.

4) The backend needs to transform the small-degree approximation polynomial on each subdomain into a polynomial with FP coefficients, suitable for evaluation in FP arithmetic, and generate code for that FP-based polynomial evaluation.

# A. Domain splitting

Given a target accuracy  $\varepsilon$  and a maximum degree d, the backend starts with computing a list of intervals  $I_k$  touching each other in so-called split-points  $s_k$ , i.e. such that  $I_{k-1} \cap I_k = \{s_k\}$ , covering the whole interval  $I = \bigcup_k I_k$  and such that for each k there exists a polynomial  $p_k \in \mathbb{R}[x]$  of degree no larger than d approximating f on  $I_k$  with a relative error at most  $\varepsilon$ :

$$\forall k, \forall x \in I_k, \quad \left| \frac{p_k(x) - f(x)}{f(x)} \right| \le \varepsilon.$$
 (9)

Such a splitting can be computed using an algorithm [18] based on bisection, interpolation of f in Chebyshev nodes and application of de la Vallée-Poussin's theorem [5]. In Frankenstein, we leverage the existing Sollya procedure guessdegree [6] to implement this step.

## B. Small degree polynomial approximation

On each subdomain  $I_k$ , the backend computes a polynomial approximation  $p_k \in \mathbb{R}[x]$  of degree at most d for the function f. The  $p_k$  are computed as *minimax* approximations with relative error, using a modified Remez-Stiefel algorithm [5]. However, these polynomials with real coefficients<sup>2</sup> are not immediately suitable to IEEE754 FP evaluation. Several problems arise.

When  $0 \notin I_k$ , in particular when  $I_k$  contains values  $x \in I_k$  that are significantly larger than 1, Horner evaluation (or any other Estrin-like evaluation) may behave badly. Roughly speaking, at each step  $q_{i+1}(x) = c_i + x \times q_i(x)$ , the evaluation error accumulated on  $q_i(x)$  will be amplified by multiplication with x when  $|x| \gg 1$ , not attenuated [19]. After splitting  $I_k$  if its radius is too large, we translate both the function f and the interval  $I_k$  by  $t_k$  to always be in the case when all  $x \in I_k - t_k$ 

stay small. In most cases, taking (a rounded value of) the midpoint of  $I_k$  is appropriate, while ensuring that the translated argument  $\xi = x - t_k$  can be computed exactly in FP arithmetic thanks to Sterbenz' lemma [29], [19]. Special care is used when *f* has a zero in  $I_k$  (see Section IV-C). Further,  $t_k$  may be optimized to take into account FP effects on the coefficients of the polynomial [19].

When  $I_k$  is small and the function f is symmetrical around some point in  $I_k$ , the minimax approximation polynomial  $p_k$ tends to reflect the symmetry: some monomials have very small coefficients compared to others. This effect may hinder FP Horner evaluation due to catastrophic cancellation but it can also be exploited to reduce the number of non-zero coefficients of the polynomial [10].

# C. Achieving relative error bounds for functions with zeros

Clearly, the ratio  $(p_k(x) - f(x))/f(x)$  can stay bounded only if f has no zero in the domain for  $x \in I_k$  or if  $p_k$  has a zero wherever f has one. Classical polynomial approximation theory either considers absolute error bounds or excludes functions with zeros. It can be extended to cover approximation with relative error of functions f with f(0) = 0 as follows: when f is known to have a zero of multiplicity m at x = 0, one computes a polynomial approximation  $q_k$  of  $x^{-m} f(x)$  and takes  $p_k(x) =$  $x^m \cdot q_k(x)$ . The only requirement is not to use Remez' original minimax algorithm but an enhanced version by Stiefel [5].

Since the backend anyway uses approximation domains  $I_k - t_k$  containing 0, the technique above already makes it possible to handle functions with a zero at some FP number. It suffices to take  $I_k$  small enough to contain a single zero of f and  $t_k$  equal to that zero in order to obtain  $f(t_k + \xi) = 0$  for  $\xi = 0$ . The value of  $t_k$ , can easily be computed with any numerical solver, such as Newton's iteration. No rigor is required in this step; if the zero is not correctly determined, relative error polynomial approximation will simply fail.

The technique no longer applies when f has a zero  $c \notin \mathbb{F}$ . However, in this case, f is actually never zero on the FP numbers it is to be evaluated at. This means that even if the ratio  $(p_k(x) - f(x))/f(x)$  is unbounded for  $x \in I_k$ , it stays bounded over  $I_k \cap \mathbb{F}$ . Provided that the procedure we use to bound the relative error only takes into account FP numbers, approximation of a function f with non-FP-representable zeros boils down to two subproblems: computing a minimax polynomial for f and translating the original function such that FP-based polynomial evaluation will exhibit bounded evaluation error.

The minimax polynomial can be computed as follows: the function g(x) = f(c + x) has an exact (FP-representable) zero at 0, and can be evaluated at any desired precision by computing *c* with a rigorous Newton algorithm. In the Sollya framework, *g* is implemented by an expression tree containing a constant function *c* whose evaluation algorithm searches the original domain  $I_k$  for a zero of *f* whenever and at whatever precision needed. This is enough for the Remez-Stiefel minimax algorithm to be applicable to *g*.

Once c itself and a polynomial p(x) approximating g(x) over  $I_k - c$  are known, we set  $t_k$  to c rounded to the nearest FP

<sup>&</sup>lt;sup>2</sup>As the underlying Sollya environment is FP-based, any polynomial we can exhibit has of course FP coefficients. However, as precision may be increased at will at this generation stage, the polynomials appear to have real coefficients.

number. The polynomial  $r(x) = p(t_k - c + x)$  then approximates  $f(t_k + x)$  over  $I_k - t_k \ge 0$ . At  $x = t_k$  (the FP value nearest to *c*), the translated argument  $\xi = x - t_k$  is exactly zero, hence FP Horner evaluation of *r* will return the constant coefficient of *r*, which is equal to (a rounded version of) f(c). For the FP *x* just after the zero,  $\xi$  will fit on a small number of bits due to cancellation in  $\xi = x - t_k$ , hence FP Horner evaluation will behave correctly, too. The accuracy of evaluation can be checked statically using Gappa as explained below.

Note, though, that the polynomial r passed to the code generation step then contains coefficients involving the Newtoniteration based representation of c. In contrast, in the usual case (when f has no zero in the domain), the coefficients directly come from the Remez-Stiefel minimax algorithm.

## D. FP code generation for polynomials

Finally, the backend needs to generate a code sequence, based on IEEE754 FP arithmetic, for each subdomain  $I_k$ , and to connect these code sequences through a branching sequence determining the appropriate subdomain given an input  $x \in I$ . Generating the subdomain determination sequence is easy.

Generating the polynomial evaluation is harder. The backend takes the following four sub-steps.

1) It determines the minimal precisions needed for the coefficients of the approximation polynomial  $p_k$  on each subdomain, following the approach described in [19].

2) It replaces the polynomial  $p_k \in \mathbb{R}[\xi]$  with a polynomial  $\tilde{p}_k \in \mathbb{R}[\xi]$  with FP coefficients. This could be done by rounding the real coefficients, however, equi-oscillation properties of the minimax polynomial and therefore accuracy might be affected too much. We therefore use a technique [5] based on lattice reduction which globally searches for a FP-valued-coefficients polynomial "close" to the original real-coefficients polynomial.

3) Starting with the target accuracy  $\varepsilon$ , the backend estimates suitable accuracies for the intermediate steps of a Horner evaluation of  $\tilde{p}_k$ . This choice of accuracies determines the FP precisions—and, possibly, double-double or triple-double expansions—to be used in the different steps. The technique is described in [10], [19]. The backend then outputs C code for polynomial evaluation along with a Gappa [8] proof script.

4) Finally, the backends runs the proof script using Gappa, hence verifying the correctness of the accuracy estimates established in the previous step.

# E. Evaluation-based vs. polynomial-based interfaces

In a cleanly designed special function code generator, the backend would take the rough polynomial approximations produced by the frontend and proceed with refining them without ever returning back to the original function f. However, this would require the frontend to take into account the relative error due to replacing f by a polynomial, which requires some understanding of FP properties of the eventual implementation when f has zeros in the domain.

For that reason and for reasons due to the reuse of existing code, Frankenstein works in a slightly different way. The backend binds a Sollya function object, corresponding to f, to the evaluator dedicated to tiny intervals provided by the frontend. This would in principle be enough for the backend to run, since the backend—in all steps, even its polynomial approximation ones—is purely based on (interval or point) evaluation. In practice, this simple interface is not enough. The number of function evaluations is too large to allow for reasonable code generation performance. In addition, an evaluation-based interface makes it hard to capture non-local properties of the function f, such as monotonicity.

We therefore enhanced Sollya to allow a function object, such as the one bound to the frontend's representation for f, to be annotated with an alternate approximate representation along with a bound on the approximation error. The approximation polynomials generated by the frontend provide such an approximate representation. When it needs to evaluate the function f at some point or over some interval, Sollya first tries to use the annotation. If this first try is inconclusive, for instance if the approximation error is too large in view of the evaluation precision, it falls back to the original evaluator bound to the frontend. Interval evaluation using the annotation exploits global properties of the polynomial. In particular, monotonicity over the whole annotation interval is detected once and for all and later used to reduce interval evaluations on subintervals to evaluations at their endpoints.

That combination has given Frankenstein the necessary performance. We should mention however that making the annotation mechanism usable for our purposes required some fine-tuning in the Sollya architecture. In particular, as we have seen, the backend manipulates not only f itself, but also expression trees containing compositions of f with other functions, which need to be reflected on any polynomial annotation. In addition, code written in the evaluation-based model tends to perform high-precision evaluations with no real need (that is, to request information on f that cannot have any influence on the correctness of the generated code). The whole benefit of the polynomial annotations can be lost if too many evaluations fall back to a slow arbitrary-precision evaluator that will try to satisfy these requests.

### V. EXAMPLES AND EXPERIMENTS

## A. Bounded intervals

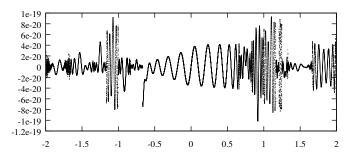
As already mentioned, there is no complete guarantee that Frankenstein succeeds to implement a given D-finite function. The following experiments show how it performs on classical special functions. All these examples were handled automatically, with minimal manual setup. The reported timings were measured on a typical desktop computer.

We start with a simple example where the functions stays away from zero on the whole implementation domain.

*Example 1:* The complementary error function  $\operatorname{erfc}(x) = 1 - 2\pi^{-1/2} \int_0^x e^{-t^2} dt$  can be defined as the solution of

$$f''(x) + 2xf'(x) = 0$$
,  $f(0) = 1$ ,  $f'(0) = -2/\sqrt{\pi}$ .

We implemented erfc in the range I = [-2; 2], with a target accuracy  $\varepsilon = 2^{-62}$ , under the constraint that no generated



0.3 0.25 0.2 0.15 0.1 0.15 0 -10 -5 0 5 10 Figure 3. V(x) for  $\sigma = 1, \lambda = \frac{1}{2}$ .

Figure 2. Relative overall error of generated implementation of erfc on [-2; 2].

polynomial have more than 14 non-zero coefficients. Such a target accuracy slightly beyond double precision is typical for first phases of correctly rounded implementations [24], [19]. Though we excluded them above, our code generator is able to handle such accuracies in simple cases, automatically introducing double-double expansions [28], [19] as necessary.

Figure 2 shows a plot of the relative error  $\tilde{f}(x)/f(x) - 1$  of the implementation. Code generation took around 780 seconds. The code generator split *I* into 16 subdomains of varying length. In the subdomain  $I_7 = [-2/3; 2/3]$ , the only nonzero coefficients of the generated polynomial are those of odd degree and that of degree 22, taking advantage [10] of an approximate symmetry around 0. Accordingly, the Horner evaluation code for  $x \in I_7$  performs evaluations in  $x^2$  in all but one steps. When executed, our implementation takes between 110 and 350 machine cycles per call, with most calls completing in 260 cycles. For comparison, a call to a typical libm exponential takes around 80 cycles.

We then consider an example with several zeros, none of which is representable in floating-point.

Example 2: The Airy function Ai satisfies

$$\begin{cases} \operatorname{Ai}''(x) - x \operatorname{Ai}(x) = 0, \\ \operatorname{Ai}(0) = 3^{-2/3} \Gamma(2/3)^{-1}, \quad \operatorname{Ai}'(0) = -3^{-1/3} \Gamma(1/3)^{-1} \end{cases}$$

We implemented Ai on I = [-4.5; 0], asking for an accuracy  $\varepsilon = 2^{-45}$ . Frankenstein takes 280 seconds to generate an implementation. It splits the domain into 10 subdomains. In two subdomains polynomials with some zero coefficients are used; the generated code precomputes  $x^2$  accordingly. It can be checked that all subdomains containing zeros of Ai(x) are translated by an amount equal to the double-precision number nearest to the zero, and an error plot confirms that the target accuracy is met even in the neighborhood of the zeros. Evaluation of the generated function takes 60 to 90 cycles, with most calls completing in 72 cycles.

Our last example directly parallels Harrison's implementation of Bessel functions for "small" arguments [14, Section 3].

*Example 3:* The Bessel functions  $J_0$  and  $Y_0$  are defined as solutions of Bessel's equation

$$xf''(x) + f'(x) + xf(x) = 0.$$
 (10)

We consider the function  $J_0$  on the interval I = [0.5; 42] with a target accuracy of  $\varepsilon = 2^{-45}$ . The immediate neighborhood of 0

is excluded because (10) is singular at x = 0 (its leading coefficient vanishes). For the same reason, initial values f(0), f'(0) do not make sense for an arbitrary solution of (10). However,  $J_0$  can still be defined as the unique solution whose value tends to 1 as  $x \rightarrow 0$ . As mentioned in Section III-A, Frankenstein's frontend supports such generalized initial conditions.

Code generation starting from this specification took about 33 minutes. The code generator splits the domain *I* into 18 subdomains. The approximation polynomials in all subdomains have non-zero coefficients. For some subdomains containing zeros of  $J'_0(x)$ , the implementer choses to store the coefficients as double-double expansions even though it rounds the final result to double precision. Evaluation of the generated implementation takes between 60 and 500 machine cycles, with most calls completing in 75 cycles.

# B. Implementation on the whole real line: an example

In some cases at least, automatically generated implementations on bounded intervals can be combined to implement a special function on the whole set of double-precision floatingpoint numbers. We illustrate how this can be done on a simple example. The manual steps we take are really an instance of a method of some generality, but we leave it to future work to handle such cases without human intervention.

The Voigt profile V(x) (Figure 3) is a probability distribution used in particular in spectrography, and whose computation has been the subject of abundant literature [27], [30]. It depends on two parameters  $\lambda, \sigma > 0$  and is defined for  $x \in \mathbb{R}$  as a convolution of a Gaussian distribution and a Cauchy distribution,

$$V(x) = \frac{1}{\sigma\sqrt{2\pi}}\frac{\lambda}{\pi}\int_{-\infty}^{+\infty}\frac{\exp\frac{-x^2}{2\sigma^2}}{(x-t)^2 + \lambda^2}dt$$

A change of variable yields the alternative expression

$$V(x) = \frac{1}{\sigma\sqrt{2\pi}} \frac{\lambda}{\pi} \int_{-\infty}^{+\infty} \frac{\exp\frac{(u-x)^2}{2\sigma^2}}{u^2 + \lambda^2} du,$$
 (11)

and it is not hard to see that (11) satisfies

$$\begin{cases} \sigma^4 V''(x) + 2\sigma^2 V'(x) + (x^2 + \lambda^2 + \sigma^2) V(x) = \frac{\lambda}{\pi}, \\ V(0) = \frac{1}{\sigma\sqrt{2\pi}} \exp\left(\frac{\lambda^2}{2\sigma^2}\right) \operatorname{erfc}\left(\frac{\lambda}{\sigma\sqrt{2}}\right), \quad V'(0) = 0. \end{cases}$$
(12)

(To remain in our general setting, a homogeneous ODE can be derived by differentiating one more time.) The Voigt profile with arbitrary parameters can be expressed in terms of the Faddeeva function, itself a renormalization of the complex error function. A general implementation of these functions, due to Johnson [17], is used for instance in the standard library of the Julia programming language. Here we are interested in implementing the function V(x) for fixed  $\lambda$  and  $\sigma$ , with an overall relative accuracy  $\varepsilon = 2^{-45}$ . In our experiment, we take  $\sigma = 1$  and  $\lambda = \frac{1}{2}$ , but the method applies verbatim to any moderate rational values.

We start with generating an implementation for  $x \in [0; 10]$ . With approximations having at most 11 nonzero coefficients, [0; 10] is split into 33 subdomains of width varying from about 0.1 around x = 5 to about 0.6 near the endpoints.

For large *x*, however, the computation of polynomial approximations starting with the initial values at 0 as described in Section III is no longer feasible. Also, it would not make any sense to split  $[10; 2^{1024}]$  into small subintervals. An obvious remedy is to consider  $f(\xi) = V(1/\xi)$  on (0, 0.1). A change of variable in (12) shows that

$$\sigma^4 \xi^6 f''(\xi) + 2\sigma^2 \xi^3 (\sigma^2 \xi^2 - 1) f'(\xi) + (\lambda^2 \xi^2 + \sigma^2 \xi^2 + 1) f(\xi) = \frac{\lambda \xi^2}{\pi}.$$

This equation has an *ir*regular singular point at 0, beyond the scope of the polynomial approximation method we discussed. Nevertheless, looking for solutions as formal power series yields a unique divergent series (compare to [13])

$$\xi^2 L(\xi) = \xi^2 + (2\sigma^2 - \lambda^2)\xi^4 + (1\sigma^4 - 10\sigma^2\lambda^2 + \lambda^4)\xi^6 + \cdots$$
(13)

whose coefficients again satisfy a simple linear recurrence. It is well-known that such formal solutions at infinity are asymptotic expansions of "true" analytic solutions and provide good approximations for large x [26]. In the present case, the solutions of the homogeneous part of the equation decrease exponentially as  $\xi \to 0$ ,  $\xi > 0$ , so all solutions of the homogeneous equation share the same asymptotic expansion (13).

With our values of  $\sigma$  and  $\lambda$ , we estimated that the polynomial  $\tilde{L}$  obtained by truncating  $L(\xi)$  to the order 90 satisfies  $|\tilde{L}(1/x) - x^2V(x)| < 2^{-65}$  for all x > 10. For the purposes of this experiment, this truncation order was found numerically. Rigorous bounds could likely be derived by the method of Olver [26, Chap. 7]. NumGfun does not support evaluations near irregular singular points, but as  $\tilde{L}(\xi)$  stays away from zero on [0, 0.1], we could directly pass it to the Frankenstein backend. The resulting code uses two polynomials of degree 11 with floating-point coefficients.

We then manually combined the two automatically generated codes into implementation that works correctly on the whole of  $\mathbb{F}$ . The Voigt function is even, so the final code starts by dropping the sign of the argument. Based on a simple comparison, it then calls either the code for V(x) around zero or computes  $\xi = 1/x$  (in double precision) and calls the code for V(1/x) around zero.

Figure 4 shows a plot of the relative error of the implementation of V. The plot covers all subdomains, both around zero and around infinity. It confirms that the accuracy target  $\varepsilon = 2^{-45}$  is satisfied. Evaluations take 48 to 180 cycles, with

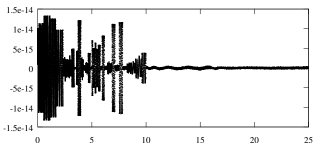


Figure 4. Relative overall error of our implementation for V(x).

most calls in the domain around zero completing in 60 cycles and most calls around infinity completing in 96 cycles.

## VI. Outlook

Our prototype code generator is a step forward towards easier, more adaptive and widespread implementation of special functions. However, it currently comes with several limitations.

First, though we expect it to succeed for all typical examples satisfying the assumptions laid out in Section II, code generation might fail on some functions. Making it work for a given function may also require some deal of human intervention.

Second, our code generator handles double precision and double/triple-double expansions, but does not work for any other FP format, let alone, say, fixed-point arithmetic. It would be interesting to extend it in that direction, perhaps with the help of other existing code generation backends.

Finally, and most importantly, automatic code generation is currently limited to bounded domains. A major perspective for future research is to provide automatic support for evaluation near infinity and singularities of the function, similar to what we outlined in the case of the Voigt profile. This also means extending our framework to handle infinite families of quasiregularly spaced zeros in the style of [14].

## ACKNOWLEDGEMENTS

The authors are grateful to S. Chevillard, F. Chyzak, B. Salvy and A. Vaugon for interesting discussions on topics related to this work, to V. Innocente and D. Piparo for suggesting the Voigt profile as a case study, and to the referees for their comments. This work was partly supported by the *Agence nationale de la recherche* grant ANR-13-INSE-0007-02 MetaLibm.

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