

The Numerical Computation of the Confluent Hypergeometric Function $U(a, b, z)$

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Summary. An algorithm is given for the computation of the confluent hypergeometric function $U(a, b, z)$. For real values of a, b and $z, z > 0$, ALGOL 60 procedures are given. The computations are based on a Miller algorithm and on asymptotic expansions.

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1. Introduction

1.1. Definitions and Relevant Properties

We consider the computation of the confluent hypergeometric function

$$U(a, b, z) = \frac{1}{\Gamma(a)} \int_0^{\infty} e^{-zt} t^{a-1} (1+t)^{b-a-1} dt. \quad (1.1)$$

This representation is valid for $\operatorname{Re} a > 0, \operatorname{Re} z > 0, b \in \mathbb{C}$. For other values of a and z we define $U(a, b, z)$ by analytic continuation. In general, i.e., for general a and b values, $U(a, b, z)$ is singular at $z=0$. It is a many-valued function with respect to z and we consider for $|\arg z| < \pi$ the principal branch, which is real (if a and b are real) for $z > 0$. With respect to a (or b) $U(a, b, z)$ is an entire function. For $a=0, -1, -2, \dots$ it can be written in terms of Laguerre polynomials

$$U(-n, \alpha + 1, z) = (-1)^n n! L_n^{(\alpha)}(z). \quad (1.2)$$

If $b-a-1=n$ ($n=0, 1, 2, \dots$) it also reduces to an elementary function. It easily follows from (1.1) that

$$U(a, a+n+1, z) = \sum_{k=0}^n (a)_k \binom{n}{k} z^{-a-k}, \quad (1.3)$$

where $(a)_k = \Gamma(a+k)/\Gamma(a)$, $k=0, 1, 2, \dots$

The function $U(a, b, z)$ is a solution of Kummer's equation

$$z w'' + (b - z) w' - a w = 0. \quad (1.4)$$

A second solution is the function (also denoted by ${}_1F_1(a, b, z)$)

$$M(a, b, z) = \sum_{n=0}^{\infty} [(a)_n / (b)_n] z^n / n! \quad (1.5)$$

which is regular for all finite z -values, although it is undefined for some pairs (a, b) .

The functions $U(a, b, z)$ and $M(a, b, z)$ satisfy recurrence relations with respect to a and b . With respect to a we have

$$f_{a-1} + (b - 2a - z)f_a + a(1 + a - b)f_{a+1} = 0 \quad (1.6)$$

which is satisfied by

$$U(a, b, z) \quad \text{and} \quad M(a, b, z) / \Gamma(1 + a - b). \quad (1.7)$$

With respect to b we have

$$(b - a - 1)f_{b-1} + (1 - b - z)f_b + z f_{b+1} = 0 \quad (1.8)$$

with solutions

$$U(a, b, z) \quad \text{and} \quad \Gamma(b - a) / \Gamma(b) M(a, b, z). \quad (1.9)$$

We use here the notation of Abramowitz and Stegun (1964, Chap. 13). More information on Kummer's function can be found in this and in many other references, for instance in Slater (1960).

For certain combinations of a and b the function $U(a, b, z)$ reduces to other special functions. We list here some examples (see Abramowitz and Stegun (1964, p. 510) for a more extensive table).

a	b	z	Relation	Function
$v + \frac{1}{2}$	$2v + 1$	$2z$	$\pi^{-\frac{1}{2}} e^z (2z)^{-v} K_v(z)$	modified Bessel
$-n$	$\alpha + 1$	z	$(-1)^n n! L_n^\alpha(z)$	Laguerre polynomial
$1 - \alpha$	$1 - \alpha$	z	$e^z \Gamma(\alpha, z)$	incomplete gamma
1	$1 + \alpha$	z	$e^z z^{-\alpha} \Gamma(\alpha, z)$	
$-\frac{1}{2}v$	$\frac{1}{2}$	$\frac{1}{2}z^2$	$2^{-\frac{1}{2}v} e^{\frac{1}{2}z^2} D_v(z)$	parabolic cylinder

The incomplete gamma function has a special cases the exponential integrals, the sine and cosine integrals and the error functions. $D_n(z)$ is an Hermite polynomial, $D_{-n-1}(z)$ is a repeated integral of the error function ($n = 0, 1, 2, \dots$).

1.2. Contents of the Paper

The algorithms given here can be used for complex values of a, b and z . In order to formulate concrete stability conditions it is better to concentrate on real a, b and $z = x > 0$.

The algorithms are implemented in two ALGOL 60 procedures:

1. the procedure *ch_u* computes the values

$$u_k = (a)_k U(a+k, b, x) \quad \text{and} \quad (a)_K U'(a+K, b, x) \quad (1.10)$$

for $a \geq 0, k=0, 1, \dots, K, b \in \mathbb{R}, x > 0$, where K is an integer ≥ 0 .

2. the procedure *uab_x* computes the values

$$U(a, b, x) \quad \text{and} \quad U'(a, b, x) \quad (1.11)$$

for $a \in \mathbb{R}, b \in \mathbb{R}, x > 0$.

The prime denotes differentiation with respect to x . The single value $(a)_K U'(a+K, b, x)$ delivered by *ch_u* is used for a backward recursion process and it plays an important role when *ch_u* calls recursively itself. Also *uab_x* calls *ch_u*, and then the derivative is important too. It is possible to obtain the derivative $U'(a+K, b, x)$ from the values u_K, u_{K-1} , for instance by using

$$x U'(a, b, x) = (a-b+x) U(a, b, x) - U(a-1, b, x).$$

For small values of x and/or large values of b this formula is not stable. Our methods in *ch_u* guarantee a stable computation for the derivative of $U(a+K, b, x)$.

In Sect. 2 we discuss recursion with respect to a . For values of x bounded away from zero we can use a Miller algorithm. For x -values close to zero, we use asymptotic expansions. In an earlier publication Temme (1975a) we used analogous methods for the computation of (1.10) with $a=b=1$.

In Sect. 3 we consider some aspects of recursion with respect to b . The ALGOL 60 procedures are described in Sect. 4. The procedures call for procedures published earlier in Temme (1975b), a publication on the computation of modified Bessel functions.

2. Recursion with Respect to a

From (1.6) and (1.7) it follows that u_k of (1.10) satisfies

$$(a+k-1)u_{k-1} + (b-2a-x-2k)u_k + (a+k+1-b)u_{k+1} = 0, \quad (2.1)$$

a second solution being

$$g_k = [\Gamma(a+k)/\Gamma(1+a+k-b)] M(a+k, b, x). \quad (2.2)$$

From asymptotic expansions of the gamma functions and from Slater (1960, p. 80) it follows easily that for $k \rightarrow \infty$

$$\begin{aligned} u_k &\sim [2/\Gamma(a)](k/x)^{(b-1)/2} e^{x/2} K_{b-1}[2(kx)^{\frac{1}{2}}][1 + O(k^{-\frac{1}{2}})], \\ g_k &= \Gamma(b)(k/x)^{(b-1)/2} e^{x/2} I_{b-1}[2(kx)^{\frac{1}{2}}][1 + O(k^{-\frac{1}{2}})], \end{aligned} \quad (2.3)$$

where $I_\nu(z)$ and $K_\nu(z)$ are modified Bessel functions. For $\text{Re } kx \rightarrow \infty$ we obtain by using well known expansions for the Bessel function

$$\begin{aligned} u_k &\sim [\pi^{\frac{1}{2}}/\Gamma(a)](k/x)^{(b-1)/2} (kx)^{-\frac{1}{2}} \exp[\frac{1}{2}x - 2(kx)^{\frac{1}{2}}], \\ g_k &\sim \frac{1}{2}\pi^{-\frac{1}{2}}\Gamma(b)(k/x)^{(b-1)/2} (kx)^{-\frac{1}{2}} \exp[\frac{1}{2}x + 2(kx)^{\frac{1}{2}}]. \end{aligned} \quad (2.4)$$

So, in the terminology of Gautschi (1967), u_k is a minimal solution and g_k is a dominant solution of (2.1). Hence the computation of u_{k+1} from u_k, u_{k-1} (using (2.1)) is not stable. If we want to use (2.1) for the computation of $\{u_k\}$ we have to use it backwards, i.e. from u_k, u_{k+1} we compute u_{k-1} . Backward recursion may be unstable for small k ; see § 3.2.

2.1. A Miller Algorithm for $\{u_k\}$

As follows from (2.4) we can use a Miller algorithm for the computation of $\{u_k\}$. For details of such an algorithm we refer to Gautschi (1967). As normalization we use

$$\sum_{k=0}^{\infty} m_k u_k = x^{-a}, \quad m_k = (-1)^k \binom{b-a-1}{k}. \quad (2.5)$$

This relation is easily verified by substituting (1.10) in (2.5) and using (1.1). For $x > 0$, $a \geq 0$, $b \in \mathbb{R}$ the u_k are non-negative; for $b \leq a+1$ all m_k are non-negative. Hence in these cases (it will appear that we can restrict b to $[0, 1]$) the series in (2.5) has non-negative terms, which is important in the numerical procedure. For $a=0$ we have $u_0=1$, $u_1=u_2=\dots=u_k=0$. For the Miller algorithm we suppose $0 < a \leq 1$, other positive a -values need not be considered.

The Miller algorithm for the recursion relation (2.1) is also considered by Wimp (1974), who in fact gives two different Miller algorithms for the computation of the U -function. His first algorithm is based on a third order difference scheme and gives a single value $U(a, b, x)$. It converges faster than his second algorithm which is similar to ours and which gives any desired number of values u_k .

Wimp did not go into the numerical details, although some important problems may arise. For instance, we found that backward recursion by using (2.1) may be unstable for certain combinations of the parameters, especially when b is large with respect to $a+k+x$. More details on this point are discussed in § 3.2.

This algorithm was also used in Temme (1975b) for the computation of the Bessel function $K_\nu(x)$. For x -values satisfying $1 < x \leq 4$, Campbell (1980) modified this algorithm by using the Wronskian relation for the modified Bessel functions. His faster method applies also in the present case. For $1.4 < x \leq 6.5$ we use (instead of (2.5)) the normalization

$$U(a, b, x) M'(a, b, x) - U'(a, b, x) M(a, b, x) = \frac{\Gamma(b)}{\Gamma(a)} x^{-b} e^x. \quad (2.6)$$

The M -functions are computed by using (1.5); the rate of convergence of (1.5) is the same as that of the exponential function ($a=b$).

In (2.6) U , M and M' are positive and U' is negative for $a > 0$, $b \geq 0$, $x > 0$. Therefore, (2.6) is stable for the indicated ranges of a , b and x . The Wronskian (2.6) is also used in the case $x > 6.5$ and $a=b$. In that case the M -functions are exponentials and (2.6) becomes

$$U(a, a, x) - U'(a, a, x) = x^{-a}.$$

For small a and/or b the gamma functions in (2.6) are not easy to handle in calculations. In the computer program we combine these functions with M' and U' in such a way that underflow or overflow will never occur when a or b is small.

In the algorithm a different version of (2.1) is used. From (2.1) it is not clear whether backward recursion is stable for small values of k . By introducing

$$v_k = (a)_k U'(a+k, b, x) \tag{2.7}$$

(compare (1.10)) we obtain the first order (2×2) -system

$$\begin{aligned} v_k &= v_{k+1} - u_{k+1} \\ u_k &= [-x v_{k+1} + (a+k+1+x-b) u_{k+1}] / (a+k) \end{aligned} \quad k \geq 0. \tag{2.8}$$

If $x > 0, a > 0, b \in \mathbb{R}$, then v_k is negative. Hence, backward recursion is stable for all $k \geq 0$ if $a+1+x-b \geq 0$. As mentioned earlier, we take $a \in (0, 1], b \in [0, 1]$ and $x > 0$. Hence (2.8) is stable for these limited ranges of a and b . Other values of b are treated in Sect. 3.

The starting value v in the Miller algorithm is computed as in Temme (1975b). In fact it is the method of Olver (1967). We take as starting values

$$\bar{u}_v = 1, \quad \bar{v}_v = -2v/[x + (x^2 + 4xv)^{\frac{1}{2}}].$$

This choice follows from approximating the differential equation for $y = U'(a, b, x)/U(a, b, x)$ viz. $x(y' + y^2) + (b-x)y - a = 0$, by putting $y' = b = 0$.

2.2. Backward Recursion with Computed Starting Values

For small values of x , the Miller algorithm is not very efficient to generate u_k of (1.10). As follows from (2.4) the dominance of g_k over u_k becomes rather weak if x is small. For small x we use computed starting values for the recursion (2.8). These values are obtained from asymptotic expansions of $U(a, b, x)$ and $U'(a, b, x)$ for large a , which are valid for small x . Such expansions are given in Slater (1960, p. 80). Similar expansions are derived in Temme (1979), where for the case of real a, b and $x > 0$ simple bounds are given for the remainders in the expansions. The expansions are for $N = 1, 2, 3, \dots$

$$\begin{aligned} U(a, b, x) &= \sum_{n=0}^{N-1} c_n(b, x) \phi_n(a, b, x) + R_N(a, b, x) \\ U'(a, b, x) &= \sum_{n=0}^{N-1} d_n(b, x) \phi_{n-1}(a, b, x) + T_N(a, b, x) \\ \phi_n(a, b, x) &= 2e^{x/2} / \Gamma(a) (x/a)^{(n+1-b)/2} K_{n+1-b} [2(ax)^{\frac{1}{2}}] \\ c_n(b, x) &= \sum_{j=0}^n b_j^{(1)}(x) b_{n-j}^{(2)}(b), \\ d_n(b, x) &= \sum_{j=0}^n b_j^{(1)}(x) b_{n-j}^{(3)}(b). \end{aligned} \tag{2.9}$$

The $b_j^{(i)}$ are defined by the generating functions (for $|\tau| < \pi$)

$$\begin{aligned} \exp[x\mu(\tau)] &= \sum_{j=0}^{\infty} b_j^{(1)}(x) \tau^j; & [\tau/(1-e^{-\tau})]^b &= \sum_{j=0}^{\infty} b_j^{(2)}(b) \tau^j; \\ b_j^{(3)}(b) &= (1-j/b) b_j^{(2)}(b); & \mu(\tau) &= 1/\tau - 1/(e^\tau - 1) - \frac{1}{2}. \end{aligned} \quad (2.10)$$

The function K_ν in ϕ_n is the modified Bessel function. Coefficients $b_j^{(1)}(x)$ and $b_j^{(2)}(b)$ are for $j=0, \dots, 8$ incorporated in the ALGOL 60 program (in fact $b_j^{(2)}(b)/b$); see arrays $\ell x, \ell b[0:8]$ in the block announced by "if $x \geq 1.4$ then".

The remainders R_N and T_N are bounded by simple expressions. Let

$$K(d, b, x) = |\sin d|^{-b} \exp[\frac{1}{2}x(1/d + 1/|\sin d|)] \quad (2.11)$$

then, as proved in Temme (1979), for $a > 0, x > 0, b \geq 0$

$$\begin{aligned} |R_N(a, b, x)| &< d^{b-N} K(d, b, x) \phi_N(a, b, x) \\ |T_N(a, b, x)| &< d^{b-N+1} K(d, b+1, x) \phi_N(a, b+1, x), \end{aligned} \quad (2.12)$$

where d is arbitrary in $[3\pi/2, 2\pi)$.

We use the above expansions for the computation of u_k and v'_k for a large value of k (see (1.10) and (2.7)). We fix N ; in the computer program we use $N=9$. Furthermore we take $0 \leq b \leq 1$. We have to choose $k \geq K$ so large that, given $a \in (0, 1], b \in [0, 1], x > 0, \varepsilon > 0, d \in [3\pi/2, 2\pi)$,

$$(a)_k |R_N(a+k, b, x)| < \varepsilon u_k \quad \text{and} \quad (a)_k |T_N(a+k, b, x)| < \varepsilon |v'_k|. \quad (2.13)$$

The minimal value of k satisfying both inequalities of (2.13) is computed by using inequalities for $\phi_j(a, b, x)$ as given in Temme (1979). Also the value of d is computed according to a device given there.

When u_k and v_k are computed we use (2.8) for backward recursion.

2.3. Negative a -Values

The integral (1.1) defines $U(a, b, z)$ for $\text{Re } a > 0$. However, as follows from recursion with respect to a , see (1.6), $U(a, b, x)$ is an entire function of a and recursion can be used for the analytic continuation of (1.1) to $\text{Re } a \leq 0$. First values for the recursion can be obtained from the algorithms of Sect. 2. We give an ALGOL 60 program *uabx* which computes $U(a, b, x)$ and the x -derivative for $a \in \mathbb{R}, b \in \mathbb{R}, x > 0$. Computation of (1.10) for $a < 0$ is not attractive due to possible singularities (for negative integer values of a) of the factor $\Gamma(a+k)/\Gamma(a)$. Special values for $a = -n, n=0, 1, 2, \dots$, follow from (1.2).

For $a \rightarrow -\infty, b$ bounded, $x > 0$ we have

$$\begin{aligned} U(a, b, x) &= \Gamma(\frac{1}{2}b - a + \frac{1}{4}) \pi^{-\frac{1}{2}} e^{\frac{1}{2}x} x^{\frac{1}{2} - \frac{1}{2}b} \cos(\chi + a\pi) [1 + O(-a)^{-\frac{1}{2}}] \\ M(a, b, x) &= \Gamma(b) \pi^{-\frac{1}{2}} e^{\frac{1}{2}x} (-ax)^{\frac{1}{2} - \frac{1}{2}b} \cos(\chi) [1 + O(-a)^{-\frac{1}{2}}] \end{aligned}$$

with $\chi = (2bx - 4abx)^{\frac{1}{2}} - \frac{1}{2}b\pi + \frac{1}{4}\pi$. It follows that $U(a, b, x)$ and $M(a, b, x)/\Gamma(1 + a - b)$, the two solutions of (1.6), are not dominant with respect to each other. Thence recursion is possible for both solutions in negative a -direction.

We use (2.8) in the form

$$\begin{aligned} U'(a, b, x) &= U'(a + 1, b, x) - U(a + 1, b, x) \\ U(a, b, x) &= -U'(a + 1, b, x) + (a + x + 1 - b)U(a + 1, b, x). \end{aligned}$$

It is not possible to give strict conditions for the stability as was done for $a > 0$. In fact $U(a, b, x)$ and $U(a + 1, b, x)$ may have different signs when $a < 0$. The same remark applies for the derivatives.

When $a < 0$ and $a = b$ we have the incomplete gamma case, viz.

$$U(-a, -a, x) = e^x \Gamma(1 + a, x).$$

In that case we can claim stability since we can recur according to the relation

$$e^x \Gamma(a + 1, x) = a e^x \Gamma(a, x) + x^a,$$

which is stable for increasing (positive) a . In terms of the U -functions the recursion is

$$U(-a, -a, x) = aU(1 - a, 1 - a, x) + xU(1 - a, 2 - a, x),$$

in which the last term is an elementary function (see (1.3)).

3. Recursion with Respect to b

For convergence aspects of the algorithms of the previous section we restricted b to the interval $[0, 1]$. Here we are concerned with the remaining b -values and we will start with $b < 0$. Throughout this section we suppose that $a > 0$ and $x > 0$.

3.1. Negative b -Values

The crucial relation is the reflection formula

$$U(a, b, x) = x^{1-b} U(1 + a - b, 2 - b, x). \tag{3.1}$$

If b is negative the b -place in the U -function on the right is positive (in fact the reflection occurs at $b = 1$, for convenience we use it at $b = 0$). In order to compute

$$\begin{aligned} u_k &= (a)_k U(a + k, b, x), \quad k = 0, \dots, K, \\ v_K &= (a)_K U'(a + K, b, x) \end{aligned} \tag{3.2}$$

for $b < 0$ we first compute

$$\begin{aligned} \bar{u}_k &= (c)_k U(c + k, 1 - b, x), \quad k = 0, \dots, K, \quad c = 1 + a - b \\ w &= (c)_K U'(c + K, 1 - b, x). \end{aligned} \tag{3.3}$$

Using (3.1) and $U'(a, b, x) = -aU(a+1, b+1, x)$, we write this as

$$\begin{aligned}\bar{u}_k &= (c)_k x^b U(a+k+1, b+1, x) = -x^b \frac{(c)_k}{(a)_{k+1}} v_k \\ w &= -(c)_{K+1} U(c+K+1, 2-b, x) = -x^{b-1} \frac{(c)_{K+1}}{(a)_{K+1}} u_{K+1}\end{aligned}\quad (3.4)$$

where $v_k = u'_k$ (see (2.7)).

For computing the values in (3.3) we need an algorithm for computing $\{u_k\}$ for $b > 1$. Details on this point will be discussed in the next subsection.

The values in (3.4) are used to compute the required u_k . First u_K is computed by using

$$aU(a, b, x) = a(1+a-b)U(a+1, b, x) - xU'(a, b, x) \quad (3.5)$$

and the remaining u_k ($k = K-1, \dots, 0$) follow from the second of (2.8). This recursion is stable since v_k and b are negative. Also (3.5) is stable.

Remark. Instead of (3.3) we might have computed

$$\begin{aligned}\bar{u}_k &= (c)_k U(c+k, 2-b, x), \quad k=0, \dots, K, \quad c=1+a-b, \\ w &= (c)_K U(c+K, 2-b, x) = -(c)_{K+1} x^{b-2} U(a+K, b-1, x).\end{aligned}$$

Then

$$\bar{u}_k = \frac{(c)_k}{(a)_k} x^{b-1} u_k;$$

hence u_k follows rather straightforwardly from the computed \bar{u}_k . However, we need also v_K . This value can be computed from the u_K , u_{K-1} , or w , u_K by using one of the recursions

$$\begin{aligned}xU'(a, b, x) &= (a-b+x)U(a, b, x) - U(a-1, b, x), \\ xU'(a, b, x) &= (1-b)U(a, b, x) - (1+a-b)U(a, b-1, x).\end{aligned}$$

For small values of x and/or large values of $-b$ they are not stable. In the approach described earlier all computations are stable.

3.2. The Case $b > 1$

Recursion with respect to b can be done by using

$$(b-a-1)U(a, b-1, x) + (1-b-x)U(a, b, x) + xU(a, b+1, x) = 0, \quad (3.6)$$

of which a second solution is $[\Gamma(b-a)/\Gamma(b)]M(a, b, x)$. From the series (1.5) it follows that $M(a, b, x) = 1 + O(b^{-1})$ for $b \rightarrow \infty$, a and x bounded. From (1.1) it follows that for $b-a-1 \geq 0$ (by using $(1+t)^{b-a-1} \geq 1$)

$$U(a, b, x) > x^{1-b} \Gamma(b-1)/\Gamma(a). \quad (3.7)$$

Hence for the recursion (3.6) the function $U(a, b, x)$ is dominant with respect to the second solution. It follows that recursion in the forward b -direction is stable.

More insight is gained when we use the derivative as in (2.8). Let us write

$$\begin{aligned} f_k &= U(a, b+k, x) \\ g_k &= U'(a, b+k, x). \end{aligned} \tag{3.8}$$

Then the recursion in the b -direction is given by

$$\begin{aligned} f_{k+1} &= f_k - g_k \\ xg_{k+1} &= (k+b)g_k - af_k. \end{aligned} \tag{3.9}$$

Since $f_k > 0$, $g_k < 0$ ($k \geq 0$), (3.9) is a recursion without subtractions (formula (3.6) lacks this property), and hence it is stable.

Suppose we want to compute (1.10) with $b > 1$. Then we define $b_1 = b - [b]$ and we compute the values

$$({}_a)_K U(a+K, b_1, x), \quad ({}_a)_K U'(a+K, b_1, x)$$

by using the algorithms of Sect. 2. Then we use (3.9) for obtaining u_K and v_K and then (2.8) can be used for the remaining u_k ($k = K-1, \dots, 0$).

As mentioned earlier, (2.8) is stable if $a+k+1+x-b$ is not negative. Large b values may violate this condition and in fact the second of (2.8) is not stable for large b -values. To show this we need the asymptotic relation (compare (3.7))

$$U(a, b, x) \sim x^{1-b} \Gamma(b-1)/\Gamma(a), \tag{3.10}$$

which is valid for $x \rightarrow 0$ ($b > 1$) or $b \rightarrow \infty$ (x fixed). Inserting this in the second of (2.8) (with modifications for v_{k+1} and u_{k+1}) we infer that indeed large values will cancel each other in order to obtain a smaller value, especially when k is small. Repeated application of (2.8) is allowed as long as k is so large that $a+k+1+x-b$ is not negative. If a, b and x (and K) are such that for some k this quantity becomes negative, instabilities may arise.

It is not easy to formulate an "if and only if" condition for the stability. The subtraction $p-q$ of two positive numbers ($q < p$) is harmless if, say, $q < \frac{1}{2}p$. The recursion (2.8) can be done by checking this criterion. However, in the computer program it is convenient to have a priori information on safe k -values. We use simply the above condition: a k -value is safe if $a+k+1+x-b$ is non-negative. Call m the largest value of k that makes $a+k+1+x-b$ negative. Then

$$\{u_{m+1}, u_{m+2}, \dots, u_K\} \tag{3.11}$$

are computed via (2.8) and for

$$\{u_0, \dots, u_m\} \tag{3.12}$$

we need a different approach.

The value m is defined by

$$m = [b-x-1-a]; \tag{3.13}$$

if $b-x-1-a=[b-x-1-a]$ then $m:=b-x-2-a$. It may happen that $m < 0$. In that case all u_k are obtained by (2.8). If $m \geq K$ none of the u_k are obtained by (2.8). In other words, the sets in (3.11) or (3.12) may be empty.

To obtain the values in (3.12) we may suppose that the values

$$(a)_j U(a+j, b_1, x), \quad j=0, \dots, m, \quad b_1 = b - [b], \quad (3.14)$$

are available, together with their derivatives. We can use (3.9) for recursion up to b with

$$f_k = (a)_j U(a+j, b_1+k, x), \quad g_k = (a)_j U'(a+j, b_1+k, x). \quad (3.15)$$

All these recursions are stable. However, it may be rather expensive, since every element in (3.14) is recurred to the b -level. For an alternative method, which may be much more efficient, we proceed as follows. We start with the element with $j=0$ in (3.14) and we recur up to $b-m$. That is, we compute (3.15) with $j=0$ for $k=0, 1, \dots, [b]-m$, giving

$$f_{[b]-m} = U(a, b-m, x), \quad g_{[b]-m} = U'(a, b-m, x).$$

Next we compute

$$F_j = (a)_j U(a+j, b-m+j, x), \quad G_j = (a)_j U'(a+j, b-m+j, x) \quad (3.16)$$

for $j=1, 2, \dots, m$. (Consider the (a, b) -plane. We compute U and U' along a diagonal in the (a, b) -plane.) These diagonal elements F_j, G_j are obtained by using

$$\begin{aligned} aU(a+1, b+1, x) &= -U'(a, b, x) \\ a x U'(a+1, b+1, x) &= -aU(a, b, x) + (b-x)U'(a, b, x), \end{aligned} \quad (3.17)$$

which in fact is Kummer's equation (1.4). In terms of F_j, G_j it reads as

$$\begin{aligned} F_{j+1} &= -G_j \\ xG_{j+1} &= -(a+j)F_j + (b+j-m-x)G_j \end{aligned} \quad (3.18)$$

for $j=0, 1, \dots, m-1$. This recursion is stable: G_j is negative and $(b+j-m-x)$ is positive for $j=0, \dots, m-1$ (cf. (3.13)). From the diagonal a final recursion in the b -direction is performed by using (3.9) with starting values F_j, G_j in order to obtain (3.12). The procedure is illustrated in Fig. 1.

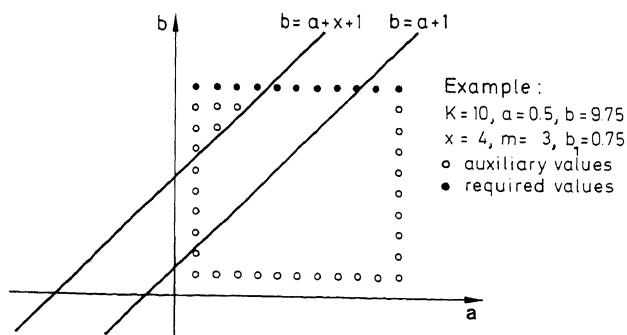


Fig. 1. The values ● at the right of the line $b = a + x + 1$ are computed via (2.8)

In the procedure *chu* this algorithm is controlled by the Boolean variables r and s ; $r = \mathbf{true}$ means that (3.11) is empty, all required values follow from diagonal elements; $s = \mathbf{false}$ means that (3.12) is empty, all required values follow from backward a -recursion. In ALGOL 60 notation: $r := K \leq m, s := m \geq 0$ (in *chu* \mathcal{K} is replaced by $kmax$).

3.3. The Polynomial Case

The actual algorithm for the case $b > 1$ is more intricate than described above. The point is that we take advantage of the possibility that some or all of the desired values $\{u_k\}$ are elementary functions. See (1.3). In that case the time-wasting algorithms of Sect. 2 may be circumvented.

Let us introduce

$$c = b - a - 1 \tag{3.19}$$

and we suppose here that c is a non-negative integer. Then the values

$$\{u_0, u_1, \dots, u_c\} \tag{3.20}$$

are polynomials, and the values

$$\{u_{c+1}, \dots, u_K\} \tag{3.21}$$

are higher transcendentals. The set in (3.21) may be empty. If it is not, its elements are computed as in the previous subsection. Since m (see (3.13)) is not larger than c , the set (3.21) is a subset of (3.11). Hence, the higher transcendentals (3.21) can be computed by backward a -recursion, whereas some of the set (3.20) (i.e., u_0, \dots, u_m) are to be computed from diagonal elements. These diagonal elements are elementary functions as well. If (3.21) is empty then all $\{u_k\}$ are polynomials. Then the first $\{u_0, \dots, u_m\}$ are computed from elementary diagonal elements, the remaining with backward a -recursion with elementary starting values.

The polynomial case is recognized in *chu* by the Boolean variables μ and q : $\mu = \mathbf{true}$ iff c is a non-negative integer, $\mu = \mathbf{true}$ and $q = \mathbf{true}$ iff all $\{u_k\}$ are polynomials. That is, if $K \leq c$. In ALGOL 60 notation: $\mu := c = \mathit{entier}(c) \wedge c \geq 0, q := \mathcal{K} \leq c$.

If we combine the possible cases of §3.2 with those of the present subsection we obtain 8 different situations A, B, \dots, H . An illustration by means of the (a, b) -plane is again very instructive. In Fig. 2 the positions of the (a, b) parameters of the desired elements $\{u_k\}$ are depicted with respect to the lines $b = a + 1, b = a + x + 1$.

For convenience, $\mu = \mathbf{true}, \mu = \mathbf{false}$, etc. are replaced by $p = 1, p = 0$, respectively. In the cases A, B, C, D, E diagonal elements must be used ($s = 1$). In A, B all elements follow from diagonal elements ($r = 1$). In C, D, E, F, G, H ($r = 0$) backward a -recursion is used for some (in F, G, H for all) elements. Values of q if $p = 0$ are not significant; r and s are not independent: $s = 0$ implies $r = 0$. G and H are treated as being the same: in G higher transcendentals are used in the backward a -recursion and the process is not terminated at the moment

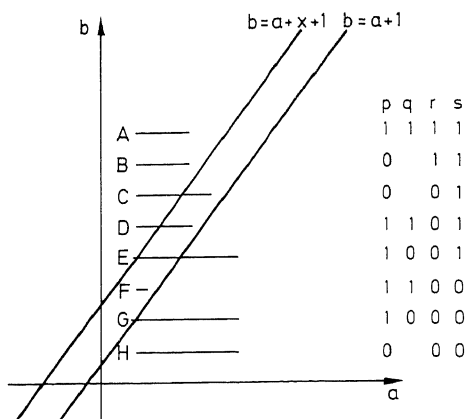


Fig. 2.

that elementary functions turn up. If $K=0$ some cases are equivalent. Then the location with respect to the line $b=a+x+1$ is not relevant; the only question is whether it is a polynomial case or not. For convenience we put $r=s=1$ (case A or B) if $K=0$.

Examples with numerical values for the different eight cases are given in Table 1.

Table 1. With the shown values of a, b, x, K all cases A through H are covered

	a	b	x	K	c	m
A	2.5	8.5	1.4	2	5	3
B	2.0	8.5	1.4	2	5.5	4
C	2.0	8.5	1.4	5	5.5	4
D	2.5	8.5	1.4	4	5	3
E	2.5	8.5	1.4	6	5	3
F	2.5	8.5	6.5	2	5	-2
G	2.5	8.5	6.5	6	5	-2
H	2.0	8.5	6.5	2	5.5	-2

4. ALGOL 60 Procedures

The procedures given here make use of external ALGOL 60 procedures for the computation of the gamma function for positive argument and of the Bessel function $K_\nu(x)$, for $x > 0$ and $\nu \in [0, 2)$. For the Bessel function we call the procedure given in Temme (1975b).

4.1. The procedure *chu*

The heading of the procedure reads as follow:

```
procedure chu(a, b, x, kmax, eps, u, uprime); value a, b, x, kmax, eps;
real a, b, x, eps, uprime; integer kmax; array u;
```

ng of the formal parameters is:

rithmetic expressions>;
 e parameters of the confluent hypergeometric function $U(a, b, x)$;
 $\geq 0, b \in \mathbb{R}, x > 0$.
 rithmetic expression>;
 e upper bound of the array u , $kmax \geq 0$.
 rithmetic expression>;
 e desired relative accuracy; $\epsilon \rho > 0$.
 rray identifier>
 rray $u[0: kmax]$;
 it: the values of $[\Gamma(a+k)/\Gamma(a)] U(a+k, b, x)$, $0 \leq k \leq kmax$, are as-
 gned to $u[k]$;
 dentifier>;
 it: the value of $[\Gamma(a+kmax)/\Gamma(a)] U'(a+kmax, b, x)$ is assigned to
prime.

ure *chu* calls for the nonlocal procedures *besska* and *gamma*; the
 ed in Temme (1975 b) (*besska* also calls for other procedures). The
 s not protected against underflow or overflow and does not give a
 ren the parameters are out of range. This last aspect is easily in-
 by the user, the first aspect strongly depends on the computing
 In general the functions $U(a+k, b, x)$ are singular at $x=0$:

- if $b < 1$, the functions are bounded at $x=0$;
- if $b = 1$, $U(a, b, x) = O(\ln x)$ as $x \rightarrow 0$;
- if $b > 1$, $U(a, b, x) = O(x^{1-b})$ as $x \rightarrow 0$.

arge b -values this singularity may cause overflow. For negative b -
 reflection formula (3.1) is used; hence, in that event, also an over-
 on may occur. For x -values bounded away from zero together with
 ies see (3.7), which shows that large function values arise in that

we have simply $u[0]=1$, $u[k]=0$, $k \geq 1$. It follows that values of
 not be obtained from

$$U(a+k, b, x) = \frac{\Gamma(a)}{\Gamma(a+k)} u[k]$$

(except for $k=0$). However, by calling the procedure with $a=1$,
chu(1, b, x, kmax, eps, u, uprime), we obtain

$$u[k] = \frac{\Gamma(1+k)}{\Gamma(1)} U(1+k, b, x), \quad k = 0, 1, \dots, kmax,$$

the values of $U(a, b, x)$ with positive integer values of a easily fol-
 at a is not restricted to $[0, 1]$.

or a are very large, underflow will occur in $u[k]$ as can be seen
 t of (2.4).

The (relative) accuracy can be controlled by *eps*. It is used to control the truncation errors in the approximation processes. Rounding errors are not considered. They may become important, although the recursions in *chu* are all strictly stable (no significant subtractions). Successive smaller choices of *eps* may yield worse results, especially if *eps* is of the same size as the machine accuracy. The relative accuracy in the external procedures *recip gamma* and *sinh* (called by *bess ka*) is about 10^{-14} .

The user may avoid a call of the Bessel function procedure *bess ka* (with its external procedures *recip gamma* and *sinh*) by skipping the part of the conditional statement announced by **if** $x \leq 1.4$ **then**. In that case the procedure will use the Miller algorithm for all $x > 0$. This will result in a less efficient algorithm with more rounding errors as x becomes smaller.

4.2. The Procedure *uabx*

For convenience we supply a function procedure which gives directly the value of $U(a, b, x)$ for $a \in \mathbb{R}$, $b \in \mathbb{R}$, $x > 0$. This procedure calls for *chu* of the previous subsection. As a second value it delivers $U'(a, b, x)$.

The heading of the procedure is:

```
real procedure uabx (a, b, x, eps, uprime); value a, b, x, eps;  
  real a, b, x, eps, uprime;
```

The meaning of the formal parameters is:

a, b, x: <arithmetic expressions>;
the parameters of $U(a, b, x)$ and $U'(a, b, x)$;
 $a \in \mathbb{R}$, $b \in \mathbb{R}$, $x > 0$.
eps: <arithmetic expression>;
the desired relative accuracy; $eps > 0$.
uprime: <identifier>;
exit: the value of $U'(a, b, x)$, the x -derivative of $U(a, b, x)$.
uabx: $uabx := U(a, b, x)$.

For underflow and overflow aspects and the role of *eps*, see the remarks in the description of *chu*.

4.3. Testing

The procedures can be compared with existing procedures for the computation of special cases of confluent hypergeometric functions. In this way we checked successfully the modified Bessel functions and the incomplete gamma functions. The procedure *chu* gives as special case the computation of the repeated integrals of the coerror function. Gautschi (1977) wrote a FORTRAN program for computing

$$i^n \operatorname{erfc} x = \int_x^\infty i^{n-1} \operatorname{erfc} t \, dt, \quad n = 1, 2, \dots$$

with $i^0 \operatorname{erfc} x = \operatorname{erfc} x$, $i^{-1} \operatorname{erfc} x = (2/\sqrt{\pi})e^{-x^2}$. In terms of the U -functions we have for $x \geq 0$

$$i^n \operatorname{erfc} x = \pi^{-\frac{1}{2}} 2^{-n} e^{-x^2} U\left(\frac{1}{2}n + \frac{1}{2}, \frac{1}{2}, x^2\right),$$

with the error function as special case for $n=0$.

The procedures can also be checked against themselves. The x -interval is divided in 3 subintervals: $(0, 1.4]$, $(1.4, 6.5]$, $(6.5, \infty)$. With

$$x_1^\pm = 1.4(1 \pm \delta), \quad x_2^\pm = 6.5(1 \pm \delta)$$

we computed $U(a, b, x_1^\pm)$ and $U(a, b, x_2^\pm)$ and we compared the results with each other. We also compared $U(a, a+n+1, x)$ with $U(a(1+\delta), a+n+1, x)$ (δ somewhat larger than the machine accuracy; the computer should recognize the polynomial case in $U(a, a+n+1, x)$ and the non-polynomial case in $U(a(1+\delta), a+n+1, x)$). All tests were satisfactory. They were done on the CD CYBER 73 of SARA, Amsterdam (machine accuracy 2^{-48}).

The boundary points 1.4 and 6.5 for the x -intervals were obtained by comparing computing time for the following choice of the parameters in $uabx$

$$a = 0.32, \quad b = 0.56, \quad eps = 10^{-10}.$$

In the immediate neighbourhood of $x=1.4$ the computing time was about 0.02 s, at $x=6.5$ it was 0.03 s. A call of chu with large values of $kmax$ and $|b|$ will require much more computing time.

Finally we give the starting index v for the Miller algorithm for $a=0.5$, $b=1$, $eps=10^{-12}$, $kmax=0$ and for several values of x :

x	1.41	2.0	6.5	6.6	10.0	50.0	100.0
v	52	40	19	38	28	12	9

4.4. Codes of chu and $uabx$

```

procedure chu( $a, b, x, kmax, eps, u, uprime$ ); value  $a, b, x, kmax, eps$ ;
real  $a, b, x, eps, uprime$ ; integer  $kmax$ ; array  $u$ ;
comment computes  $\gamma(a+k)/\gamma(a) \times u(a+k, b, x)$ 
for  $k=0$  (1)  $kmax$ , and the  $x$ -derivative of  $u(a+kmax, b, x)$ ;
if  $a < 0$  or  $x < 0$  or  $kmax < 0$  or  $eps \leq 0$  then
begin comment here the user can incorporate an output statement
with the message "parameters out of range";
end else
if  $a=0$  then
begin  $u[0] := 1$ ;  $uprime := 0$ ;
for  $kmax := kmax$  step  $-1$  until  $1$  do  $u[kmax] := 0$ 
end else
if  $b < 0$  then
    
```

```

begin real c, d, e, w; integer j; array v[0:kmax];
  c:=a-b+1; d:=x↑(-b); chu(c, 1-b, x, kmax, eps, v, w);
  for j:=0 step 1 until kmax do
    begin e:=(a+j)×d; v[j]:=-e×v[j]; d:=e/(c+j) end;
  uprime:=v[kmax]; u[kmax]:=-x×(uprime+e×w)/(a+kmax);
  c:=c+x; for j:=kmax-1 step -1 until 0 do
    u[j]:=(-x×v[j+1]+(c+j)×u[j+1])/(a+j)
  end else
  if b>1 then
  begin real a1, b1, c, d, e, f, g, h, u3, v, w; integer i, j, k, m, n; boolean p, q, r, s;
    procedure brec(a, b, k, f, g); value a, b, k; real a, b, f, g; integer k;
      begin k:=k-1; for i:=0 step 1 until k do
        begin h:=f-g; g:=((i+b)×g-a×f)/x; f:=h end
      end brec;
    n:=entier(b); b1:=b-n; a1:=a+kmax; c:=b-a-1;
    e:=c-x; m:=entier(e); if m=e then m:=m-1;
    p:=c=entier(c) ∧ c≥0; q:=kmax≤c; r:=kmax≤m; s:=m≥0;
    if kmax=0 then r:=s:=true;
    if r then m:=kmax; k:= (if p then c else n)-m;
    if ¬r then
      begin if p and q then
        begin g:=1; i:=kmax-1; for j:=0 step 1 until i do
          g:=g×(j+a); f:=g×x↑(-a1); g:=-a1×f/x;
          brec(a1, a1+1, c-kmax, f, g)
        end else
          begin chu(a, b1, x, kmax, eps, u, u3); f:=u[kmax]; g:=u3;
            brec(a1, b1, n, f, g); if ¬p and s then
              begin for j:=kmax step -1 until 1 do
                u3:=u3-u[j]; v:=u[0]; w:=u3; d:=b1
              end
            end; n:=m+1; u[kmax]:=f; uprime:=g; if ¬s then n:=0;
            for j:=kmax-1 step -1 until n do
              begin h:=(-x×g+(j-e)×f)/(a+j); g:=g-f; f:=u[j]:=h end
            end; if s then
              begin if p then
                begin v:=x↑(-a); w:=-a×v/x; d:=a+1 end else
                  if r then
                    begin chu(a, b1, x, 0, eps, u, w); v:=u[0]; d:=b1 end;
                    brec(a, d, k, v, w); e:=b-n-x;
                    for j:=0 step 1 until m do
                      begin if j=0 then
                        begin f:=u[0]:=v; g:=w end else
                          begin h:=-w; g:=w=-((a+j-1)×v-(j+e)×w)/x; v:=u[j]:=h
                        end; brec(a+j, b+j-m, m-j, u[j], g)
                      end; if m=kmax then uprime:=g
                    end
                end
              end
            end
          end
        end
      end
    end
  end
end

```



```

else
: ≤ 1.4 then
in real d, delta, e, f, p, q, r, s, t, t0, t1, u0, u1, u2, u3, v, w, x2, y, z;
integer n, nu, k0, k1, i, j; array bb, bb1, bx[0:8], fi[-1:8];
n:=9;
v:=12.56637; r:=(x-v*(b+1))/2; d:=(v*n+r-sqrt(r*r+4*x*n*x))/(2*x);
if d < 4.7124 then
begin d:=4.7124; v:=w:=1 end else
begin v:=abs(sin(d)); w:=v^(-1-b) end;
w:=w*exp(0.5*x*(1/v+1/d));
delta:=eps*exp(-0.5*x+(n-1-b)*ln(d))/w;
z:=0.5/delta; v:=0.5-b; i:=n-1;
for j:=1 step 1 until i do z:=z*(j+v); i:=0;
t:=sqrt(x)*z^(1/(2*n)); e:=ln(delta)+n-n*ln(x);
r:=n+b+t; s:=1+b+t; p:=ln(r); q:=ln(s);
f:=(ln(t+0.5)-2*x*n*ln(t)+(r-0.5)*p-(s-0.5)*q-e)/
(1/(t+0.5)-2*x*n/t+0.5*(n-1)/(r*s)+p-q);
if f < 0 then
begin t:=t-f; t:=sqrt(2*x+t*t); i:=i+1;
if i < 10 then goto lab
end else k0:=1+entier(t*t/x-a);
nu:= if kmax ≥ k0 then 1+kmax else k0;
r:=a+nu; w:=sqrt(x/r); v:=2*x*r*w; bess ka(-b, v, t0, t1);
v:=w^(-b); bb[0]:=bb1[0]:=bx[0]:=1;
a1:=fi[-1]:=v*t0; u0:=fi[0]:=v*w*t1; x2:=x*x;

x[1]:=-x/12;
x[2]:=x2/288;
x[3]:=-x*(5*x2-72)/51840;
x[4]:=x2*(5*x2-288)/2488320;
x[5]:=-x*(x2*(7*x2-1008)+6912)/209018880;
x[6]:=x2*(x2*(35*x2-10080)+279936)/75246796800;
x[7]:=-x*(x2*(x2*(x2*5-2520)+176256)-746496)/902961561600;
x[8]:=x2*(x2*(x2*(x2*5-4032)+566784)-9953280)/86684309913600;

b[1]:=0.5;
b[2]:=(3*b-1)/24;
b[3]:=b*(b-1)/48;
b[4]:=(b*(b*(b*15-30)+5)+2)/5760;
b[5]:=b*(b*(b*(b*3-10)+5)+2)/11520;
b[6]:=(b*(b*(b*(b*(b*63-315)+315)+91)-42)-16)/2903040;
b[7]:=b*(b*(b*(b*(b*(b*9-63)+105)+7)-42)-16)/5806080;
b[8]:=(b*(b*(b*(b*(b*(b*135-1260)+3150)-840)-2345)-
540)+404)+144)/1393459200;

```

```

for  $i:=1$  step 1 until  $n-1$  do
begin
   $t0:=bb[i]; t1:=bb1[i]:=(b-i) \times t0;$ 
  for  $j:=1$  step 1 until  $i-1$  do
  begin
     $t0:=t0+bb[i-j] \times bx[j];$ 
     $t1:=t1+bb1[i-j] \times bx[j]$ 
  end;
   $t0:=bx[i]+b \times t0; t1:=t1+bx[i];$ 
   $fi[i]:=(x \times fi[i-2]+(i-b) \times fi[i-1])/r;$ 
   $u0:=u0+t0 \times fi[i]; u1:=u1+t1 \times fi[i-1]$ 
end;
   $w:=2 \times \exp(x/2)/\text{gamma}(1+a);$ 
   $u2:=w \times u0; u3:=-w \times u1; v:=a+1-b+x; k1:=nu-1;$ 
  for  $j:=k1$  step -1 until 1 do
  begin  $u1:=(-x \times u3+(v+j) \times u2)/(a+j);$ 
     $u3:=u3-u2; u2:=u1; \text{if } j \leq kmax \text{ then } u[j]:=a \times u2;$ 
    if  $j=kmax$  then  $u\prime:=a \times u3$ 
  end;  $u[0]:=-x \times u3+v \times u2;$ 
  if  $kmax=0$  then  $u\prime:=a \times (u3-u2)$ 
end else
begin real  $ar, br, cr, c, er, m0, m1, mr, p0, p1, p2, q, u1, u2, u3, v, w;$ 
  integer  $k, n, r;$  boolean  $large\ x;$ 
  procedure recursion;
  begin  $p2:=(br \times p1-ar \times p0)/cr; er:=er \times ar/cr;$ 
     $r:=r+1; \text{if } large\ x \text{ then } mr:=mr \times (1+c/r); v:=er/p2;$ 
     $br:=br+2; cr:=cr+1; p0:=p1; p1:=p2$ 
  end recursion;
   $n:=\text{entier}(a); \text{if } a=n \text{ then } n:=n-1; a:=a-n; kmax:=kmax+n;$ 
   $large\ x:=x > 6.5 \wedge a \neq b;$ 
  if  $large\ x$  then  $mr:=1$  else
  begin  $mr:=0; \text{if } a=b \text{ then}$ 
  begin  $m0:=a; m1:=1$  end else
  begin  $m0:=0; m1:=v:=1;$ 
  for  $r:=1, r+1$  while  $v \geq m1 \times eps$  do
  begin  $v:=v \times v/r; m0:=m0+v; v:=v \times (a+r)/(b+r); m1:=m1+v$  end;
   $v:=\exp(-x) \times \text{gamma}(a+1)/\text{gamma}(b+1);$ 
   $m0:=v \times (b+a \times m0); m1:=v \times m1$ 
  end
  end;
   $c:=a-b; cr:=2+c; br:=x+a+cr; p0:=0; v:=p1:=er:=1; r:=0;$ 
  for  $ar:=a+r$  while  $r \leq kmax$  do recursion;  $w:=p0 \times p1/er;$ 
  for  $ar:=a+r, ar+1$  while  $v \times (w/p0+mr \times (2+a/r)) \geq eps$  do recursion;
   $c:=1+c; v:=x+c; u2:=1; w:=0; u3:=-2 \times r/(x+\text{sqrt}(x \times (x+4 \times r)));$ 
  for  $r:=r-1, r-1$  while  $r > 0$  do

```

```

begin if large x then
  begin  $w := w + mr \times u2$ ;  $mr := mr \times (r + 1)/(c + r)$  end;
   $u1 := (-x \times u3 + (v + r) \times u2)/(a + r)$ ;  $u3 := u3 - u2$ ;  $u2 := u1$ ;
  if  $r \geq n$  and  $r \leq kmax$  then  $u[r-n] := u2$ ;
  if  $r = kmax$  then  $u_{prime} := u3$ 
end;
 $u1 := -x \times u3 + v \times u2$ ;  $u3 := u3 - u2$ ;  $v := a$ ;  $k := n - 1$ ;
if  $kmax = 0$  then  $u_{prime} := u3$ ;  $kmax := kmax - n$ ;
 $w :=$  if large x then  $x \uparrow (-a)/(a \times (w + c \times u2) + u1)$  else
   $x \uparrow (-b)/(u1 \times m1 - u3 \times m0)$ ;
for  $r := 0$  step 1 until  $k$  do  $v := v/(a + r)$ ;
if  $n = 0$  then begin  $k := 1$ ;  $u[0] := w \times u1$  end else  $k := 0$ ;
 $w := v \times w$ ;  $u_{prime} := w \times u_{prime}$ ;
for  $r := k$  step 1 until  $kmax$  do  $u[r] := w \times u[r]$ 
end chu;
real procedure  $uabx(a, b, x, eps, u_{prime})$ ; value  $a, b, x, eps$ ;
real  $a, b, x, eps, u_{prime}$ ;
begin real  $a1, c, p, q, r$ ; integer  $j, n$ ; array  $u[0:0]$ ;
 $n :=$  if  $a < 0$  then  $entier(a)$  else 0;  $q := a1 := a - n$ ;  $u[0] := 1$ ;
if  $n < 0 \wedge a = b$  then
  begin if  $a1 > 0$  then  $chu(a1, a1, x, 0, eps, u, q)$ ;
     $p := u[0]$ ;  $r := p - q$ ;
    for  $j := 1$  step 1 until  $-n$  do
      begin  $r := x \times r$ ;  $q := (a1 - j) \times p$ ;  $p := r - q$  end
    end else
    begin if  $a1 > 0$  then  $chu(a1, b, x, 0, eps, u, q)$ ;
       $c := 1 + a1 - b + x$ ;  $a1 := a1 - 1$ ;  $p := u[0]$ ;
      for  $j := 1$  step 1 until  $-n$  do
        begin  $r := (c - j) \times p - x \times q$ ;  $q := (a1 - j) \times (q - p)$ ;  $p := r$  end;
      end;  $uabx := p$ ;  $u_{prime} := q$ 
end uabx;

```

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