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> DGIG_help(DGIGpdf)
*****
* ==> DGIGpdf(r1,r2,l1,l2,z,prec,prec) <==
* Computes values for the PDF of the DGIG distribution
* Its has 7 arguments, the first 5 are mandatory:
* r1 - list of shape parameters for the Gamma distributions with positive sign
* r2 - list of shape parameters for the Gamma distributions with negative sign
* l1 - list of rate parameters for the Gamma distributions with positive sign
* l2 - list of rate parameters for the Gamma distributions with negative sign
* z - running value at which the CDF is to be computed
* prec - (optional argument) number of precision digits for the computations
* (default value: 50)
* precp - (optional argument) number of digits for printing the result
* (default value: 20)
*****
> DGIGpdf(c(5,3,4),c(2,3),c(3.2,4.5,1.6),c(1.4,1.3),1.2)
0.19089711134150040569
> # One should be careful and know that for a given value of 'prec', not all 'prec' digits in
> # the result will be exact
> DGIGpdf(c(5,3,4),c(2,3),c(3.2,4.5,1.6),c(1.4,1.3),1.2,,50)
0.19089711134150040569284558610654807001361124302019
> DGIGpdf(c(5,3,4),c(2,3),c(3.2,4.5,1.6),c(1.4,1.3),1.2,70,50)
0.19089711134150040569284558610654807001361129292395
> # All user functions, besides giving a printed result, allow for saving the result in a
variable
> # which will then have all 'prec' digits
> val<-DGIGpdf(c(5,3,4),c(2,3),c(3.2,4.5,1.6),c(1.4,1.3),1.2)
0.19089711134150040569
> val
1 'mpfr' number of precision 166 bits
[1] 0.190897111341500405692845586106548070013611243020194
> val<-DGIGpdf(c(5,3,4),c(2,3),c(3.2,4.5,1.6),c(1.4,1.3),1.2,70)
0.19089711134150040569
> val
1 'mpfr' number of precision 232 bits
[1] 0.19089711134150040569284558610654807001361129292394637869101812547108985
>
> DGIG_help(DGIGcdf)
*****
* ==> DGIGcdf(r1,r2,l1,l2,z,prec,prec) <==
* Computes values for the CDF of the DGIG distribution
* Its has 7 arguments, the first 5 are mandatory:
* r1 - list of shape parameters for the Gamma distributions with positive sign
* r2 - list of shape parameters for the Gamma distributions with negative sign
* l1 - list of rate parameters for the Gamma distributions with positive sign
* l2 - list of rate parameters for the Gamma distributions with negative sign
* z - running value at which the CDF is to be computed
* prec - (optional argument) number of precision digits for the computations
* (default value: 50)
* precp - (optional argument) number of digits for printing the result
* (default value: 20)
*****
> DGIGcdf(c(5,3,4),c(2,3),c(3.2,4.5,1.6),c(1.4,1.3),1.2)
0.52826935818589085876
> val<-DGIGcdf(c(5,3,4),c(2,3),c(3.2,4.5,1.6),c(1.4,1.3),1.2)
0.52826935818589085876
> val
1 'mpfr' number of precision 166 bits
[1] 0.528269358185890858759736842929349108110114814275989
> val<-DGIGcdf(c(5,3,4),c(2,3),c(3.2,4.5,1.6),c(1.4,1.3),1.2,70,30)
0.528269358185890858759736842929
> val
1 'mpfr' number of precision 232 bits
[1] 0.52826935818589085875973684292934910811011702394545941854056503024295477
>
> DGIG_help(plotDGIGpdf)
*****
* ==> plotDGIGpdf(r1,r2,l1,l2,ul,step,prec) <==
* Plots pdfs of the DGIG distribution
* arguments are (first 5 are mandatory):
* r1 - list of shape parameters for the 'positive' part of the DGIG distribution
* r2 - list of shape parameters for the 'negative' part of the DGIG distribution
* l1 - list of rate parameters for the 'positive' part of the DGIG distribution

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[illegible]

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> DGIGcdf(c(4, 2, 3), c(2, 1), c(3.6, 2.5, 4.2), c(1.4, 5.3), q05, 100, 80)
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[illegible]