## Table 1

– Valid values of &retmode and &rettype for EFetch (null = empty string)

Record Type	&rettype	&retmode	
All Database	S		
Document summary	docsum	xml, default	
List of UIDs in XML	uilist	xml	
List of UIDs in plain text	uilist	text	
db = bioproject			
Full record XML	xml, default	xml, default	
db = biosamp	le		
Full record XML	full, default	xml, default	
Full record text	full, default	text	
db = biosystems			
Full record XML	xml, default	xml, default	
db = gds			
Summary	summary, default	text, default	
db = gene			
text ASN.1	null	asn.1, default	
XML	null	xml	
Gene table	gene_table	text	
db = homologene			
text ASN.1	null	asn.1, default	
XML	null	xml	
Alignment scores	alignmentscores	text	
FASTA	fasta	text	
HomoloGene	homologene	text	
db = mesh			
Full record	full, default	text, default	
db = nlmcatalog			
Full record	null	text, default	
XML	null	xml	
db = nuccore, nucest, nucgss, protein or popset			
text ASN.1	null	text, default	
binary ASN.1	null	asn.1	
Full record in XML	native	xml	
Accession number(s)	acc	text	

4/2019 lable 1, [- valid values of &fetimode a	and wrettype for Erec	.cii (iiuii – eilipt)		
FASTA	fasta	text		
TinySeq XML	fasta	xml		
SeqID string	seqid	text		
Additional options for db = nuccore, nucest, nucgss or popset				
GenBank flat file	gb	text		
GBSeq XML	gb	xml		
INSDSeq XML	gbc	xml		
Additional option for db = nuccore and protein				
Feature table	ft	text		
Additional option for d	b = nuccore			
GenBank flat file with full sequence (contigs)	gbwithparts	text		
CDS nucleotide FASTA	fasta_cds_na	text		
CDS protein FASTA	fasta_cds_aa	text		
Additional option for db = nucest				
EST report	est	text		
Additional option for db = nucgss				
GSS report	gss	text		
Additional options for db = protein				
GenPept flat file	gp	text		
GBSeq XML	gp	xml		
INSDSeq XML	gpc	xml		
Identical Protein XML	ipg	xml		
db = pmc				
XML	null	xml, default		
MEDLINE	medline	text		
db = pubmed	d			
text ASN.1	null	asn.1, default		
XML	null	xml		
MEDLINE	medline	text		
PMID list	uilist	text		
Abstract	abstract	text		
db = sequences				
text ASN.1	null	text, default		
Accession number(s)	acc	text		
FASTA	fasta	text		
SeqID string	seqid	text		
db = snp				
text ASN.1	null	asn.1, default		

XML	null	xml	
Flat file	flt	text	
FASTA	fasta	text	
RS Cluster report	rsr	text	
SS Exemplar list	ssexemplar	text	
Chromosome report	chr	text	
Summary	docset	text	
UID list	uilist	text or xml	
db = sra			
XML	full, default	xml, default	
db = taxonomy			
XML	null	xml, default	
TaxID list	uilist	text or xml	
db = clinvar			
ClinVar Set	clinvarset	xml, default	
UID list	uilist	text or xml	
db = gtr			
GTR Test Report	gtracc	xml, default	

From: The E-utilities In-Depth: Parameters, Syntax and More



Entrez Programming Utilities Help [Internet].

Bethesda (MD): National Center for Biotechnology Information (US); 2010-.

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