Anexos

Informações Genbank do genoma do S. aureus N315

Id: BA000018.3

Name: BA000018

Description: Staphylococcus aureus subsp. aureus N315 DNA, complete genome

Annotations:

'molecule_type': 'DNA' 'topology': 'circular',

• 'data file division': 'BCT',

'date': '07-OCT-2016',

'accessions': ['BA000018', 'AP003129-AP003138'],

'sequence_version': 3,

'keywords': ["],

'source': 'Staphylococcus aureus subsp. aureus N315',

'organism': 'Staphylococcus aureus subsp. aureus N315',

- 'taxonomy': ['Bacteria', 'Firmicutes', 'Bacilli', 'Bacillales', 'Staphylococcaceae', 'Staphylococcus'],
- 'references': [Reference(title='Whole genome sequencing of meticillin-resistant Staphylococcus aureus', ...), Reference(title='Direct Submission', ...)],
- 'comment': 'On or before Nov 5, 2004 this sequence version replaced AP003129.2,\nAP003130.2, AP003131.2, AP003132.2, AP003133.2, AP003134.2,\nAP003135.2, AP003136.2, AP003137.2, AP003138.2.'

DBXrefs: ['BioProject:PRJNA264', 'BioSample:SAMD00061099']

Genes essenciais (pelo modelo do Optflux) sem homologia no genoma humano (OrderedLocusNames e respetivo e-value)

. •			
[[['SA0016', 0.253085],	['SA0997', 0.638148],	['SA0843', 0.240757],
['SA0176', 0.174251],	['SA1104', 0.205002],	['SA0910', 0.0793319],
['SA0177', 0.0700738],	['SA1115', 0.0699459],	['SA0911', 0.808787],
['SA0178', 0.0772464],	['SA1164', 0.085164],	['SA0916', 0.429948],
['SA0179', 0.309651],	['SA1165', 0.245048],	['SA0919', 0.247518],
['SA0344', 0.570958],	['SA1177', 1.87107],	['SA0920', 0.0971463],
['SA0345', 0.123391],	['SA1197', 0.289576],	['SA0921', 0.112943],
['SA0346', 0.0883758],	['SA1199', 0.107574],	['SA0923', 0.252724],
['SA0419', 0.0869642],	['SA1201', 0.23003],	['SA0924', 0.955198],
['SA0439', 0.10084],	['SA1202', 0.715529],	['SA0925', 0.102962],
['SA0457', 0.267056],	['SA1203', 0.163862]]	['SA0926', 0.145853],
['SA0473', 0.0911547],	[['SA0016', 0.281624],	['SA0937', 0.0758166],
['SA0474', 0.425756],	['SA0134', 0.0545394],	['SA0965', 1.24871],
['SA0512', 0.285471],	['SA0176', 0.901747],	['SA0994', 0.588693],
['SA0547', 0.769982],	['SA0178', 0.626128],	['SA0996', 0.274383],
['SA0593', 0.733756],	['SA0179', 0.0531147],	['SA1065', 0.588667],
['SA0594', 0.219757],	['SA0344', 0.0599982],	['SA1088', 0.0849591],
['SA0596', 0.0753623],	['SA0345', 0.160552],	['SA1104', 1.25614],
['SA0669', 0.08767],	['SA0346', 0.372101],	['SA1115', 0.30158],
['SA0670', 0.543606],	['SA0375', 0.277368],	['SA1126', 0.391652],
['SA0683', 0.128109],	['SA0376', 1.02961],	['SA1150', 0.539172],
['SA0693', 0.740484],	['SA0439', 0.0659293],	['SA1165', 0.280725],
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['SA0794', 0.326249],	['SA0487', 0.350747],	['SA1197', 0.558275],
['SA0795', 0.0533705],	['SA0547', 0.0598298],	['SA1199', 0.454343],
['SA0842', 0.0526714],	['SA0548', 0.0506916],	['SA1201', 0.512763],
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['SA0910', 0.215554],	['SA0592', 0.13308],	['SA1204', 0.281624]
['SA0911', 0.116035],	['SA0593', 1.18195],	['SA1205', 0.0545394]
['SA0913', 0.254344],	['SA0594', 0.889915],	['SA1226', 0.901747]
['SA0916', 0.408854],	['SA0597', 0.244495],	['SA1228', 0.626128]
['SA0919', 0.0633448],	['SA0669', 0.359165],	['SA1229', 0.0531147]
['SA0920', 0.611704],	['SA0670', 0.0679558],	['SA1245', 0.0599982]
['SA0924', 0.14368],	['SA0793', 0.220178],	['SA1250', 0.160552]
['SA0938', 0.067339],	['SA0794', 0.0759321],	['SA1259', 0.372101]
['SA0965', 0.0511052],	['SA0795', 1.44111],	['SA1298', 0.277368]
['SA0996', 0.747054],	['SA0796', 0.312388],	['SA1299', 1.02961]

['SA1306', 0.0659293]	['SA1650', 0.0793319]	['SA2341', 1.25614]
['SA1346', 0.0967526]	['SA1651', 0.808787]	['SA2347', 0.30158]
['SA1397', 0.350747]	['SA1728', 0.429948]	['SA2406', 0.391652]
['SA1424', 0.0598298]	['SA1735', 0.247518]	['SA2412', 0.539172]
['SA1427', 0.0506916]	['SA1749', 0.0971463]	['SA2456', 0.280725]
['SA1439', 0.140952]	['SA1858', 0.112943]	['SA2465', 0.603203]
['SA1461', 0.13308]	['SA1861', 0.252724]	['SA2466', 0.558275]
['SA1487', 1.18195]	['SA1863', 0.955198]	['SA2467', 0.454343]
['SA1492', 0.889915]	['SA1864', 0.102962]	['SA2468', 0.512763]
['SA1494', 0.244495]	['SA1865', 0.145853]	['SA2471', 0.138754]
['SA1496', 0.359165]	['SA1938', 0.0758166]	
['SA1522', 0.0679558]	['SA1965', 1.24871]	
['SA1571', 0.220178]	['SA2027', 0.588693]	
['SA1585', 0.0759321]	['SA2136', 0.274383]	
['SA1587', 0.312388]	['SA2288', 0.588667]	
['SA1589', 0.240757]	['SA2333', 0.0849591]	

Genes essenciais tanto para o Optflux e para o DEG Sem Homologia com o genoma Humano

'SA0179', 0.309651	'SA1177', 1.87107	'SA1728', 0.429948
'SA0457', 0.267056	'SA1204', 0.281624	'SA2027', 0.588693
'SA0924', 0.14368	'SA1259', 0.372101	'SA2406', 0.391652
'SA0997', 0.638148	'SA1492', 0.889915	'SA1522', 0.0679558
'SA1104', 0.205002	'SA1494', 0.244495	'SA1346', 0.096752

Dihidrofolato redutase - UniProt

number: 1

accessions: ['P99079', 'P10167'] annotation_update: ('05-DEC-2018', 89) comments: ['FUNCTION: Key enzyme in folate metabolism. Catalyzes an essential reaction for de novo glycine and purine synthesis, and for DNA precursor synthesis (By similarity). {ECO:0000250}.', 'CATALYTIC ACTIVITY: Reaction=(6S)-5,6,7,8tetrahydrofolate + NADP(+) = 7,8- dihydrofolate + H(+) + NADPH; Xref=Rhea:RHEA:15009, ChEBI:CHEBI:15378, Chebi:Chebi:57451, Chebi:Chebi:57453, Chebi:Chebi:57783, Chebi:Chebi:58349; EC=1.5.1.3; Evidence={ECO:0000255 | PROSITE-ProRule:PRU00660};', 'PATHWAY: Cofactor biosynthesis; tetrahydrofolate biosynthesis; 5,6,7,8tetrahydrofolate from 7,8-dihydrofolate: step 1/1.', 'SIMILARITY: Belongs to the dihydrofolate reductase family. {ECO:0000305}.'] created: ('01-MAR-2005', 0) cross_references: [('EMBL', 'BA000018', 'BAB42519.1', '-', 'Genomic_DNA'), ('RefSeq', 'WP_000175746.1', 'NC_002745.2'), ('ProteinModelPortal', 'P99079', '-'), ('SMR', 'P99079', '-'), ('SWISS-2DPAGE', 'P99079', '-'), ('EnsemblBacteria', 'BAB42519', BAB42519', 'BAB42519'), ('KEGG', 'sau:SA1259', '-'), ('HOGENOM', 'HOG000040233', '-'), ('KO', 'K00287', '-'), ('OMA', 'RDNQLPW', '-'), ('BioCyc', 'SAUR158879:G1G21-1440-MONOMER', '-'), ('UniPathway', 'UPA00077', 'UER00158'), ('Proteomes', 'UP000000751', 'Chromosome'), ('GO', 'GO:0004146', 'F:dihydrofolate reductase activity', 'IEA:UniProtKB-EC'), ('GO', 'GO:0050661', 'F:NADP binding', 'IEA:InterPro'), ('GO', 'GO:0006545', 'P:glycine biosynthetic process', 'IEA:InterPro'), ('GO', 'GO:0006730', 'P:one-carbon metabolic process', 'IEA:UniProtKB-KW'), ('GO', 'GO:0046654', 'P:tetrahydrofolate biosynthetic process', 'IEA:UniProtKB-UniPathway'), ('CDD', 'cd00209', 'DHFR', '1'), ('Gene3D', '3.40.430.10', '-', '1'), ('InterPro', 'IPR012259', 'DHFR'), ('InterPro', 'IPR024072', 'DHFR-like_dom_sf'), ('InterPro', 'IPR017925', 'DHFR_CS'), ('InterPro', 'IPR001796', 'DHFR_dom'), ('PANTHER', 'PTHR22778:SF16', 'PTHR22778:SF16', '1'), ('Pfam', 'PF00186', 'DHFR_1', '1'), ('PIRSF', 'PIRSF000194', 'DHFR', '1'), ('SUPFAM', 'SSF53597', 'SSF53597', '1'), ('PROSITE', 'PS00075', 'DHFR_1', '1'), ('PROSITE', 'PS51330', 'DHFR_2', '1')] data class: Reviewed description: RecName: Full=Dihydrofolate reductase; Short=DHFR; EC=1.5.1.3; entry_name : DYR_STAAN features: [('INIT_MET', 1, 1, 'Removed. {ECO:0000250}.', "), ('CHAIN', 2, 159, 'Dihydrofolate reductase.', 'PRO_0000186408'), ('DOMAIN', 2, 157, 'DHFR. {ECO:0000255|PROSITE- ProRule:PRU00660}.', "), ('NP_BIND', 7, 8, 'NADP. {ECO:0000250}.', "), ('NP_BIND', 15, 20, 'NADP. {ECO:0000250}.', "), ('NP_BIND', 44, 47, 'NADP. {ECO:0000250}.', "), ('NP_BIND', 63, 66, 'NADP. {ECO:0000250}.', "), ('NP BIND', 93, 98, 'NADP. {ECO:0000250}.', "), ('REGION', 6, 8, 'Substrate binding. {ECO:0000250}.', "), ('BINDING', 28, 28, 'Substrate. {ECO:0000250}.', "), ('BINDING', 58, 58, 'Substrate. {ECO:0000250}.', "), ('BINDING', 112, 112, 'Substrate. {ECO:0000250}.', ")] gene_name: Name=folA; OrderedLocusNames=SA1259; host organism:[] host taxonomy id:[] keywords: ['Complete proteome', 'NADP', 'One-carbon metabolism', 'Oxidoreductase'] molecule_type: None organelle: organism: Staphylococcus aureus (strain N315). organism_classification: ['Bacteria', 'Firmicutes', 'Bacilli', 'Bacillales', 'Staphylococcaceae', 'Staphylococcus'] protein_existence: 1 references: Authors: Kuroda M., Ohta T., Uchiyama I., Baba T., Yuzawa H., Kobayashi I., Cui L., Oguchi A., Aoki K., Nagai Y., Lian J.-Q., Ito T., Kanamori M., Matsumaru H., Maruyama A., Murakami H., Hosoyama A., Mizutani-Ui Y., Takahashi N.K., Sawano T., Inoue R., Kaito C., Sekimizu K., Hirakawa H., Kuhara S., Goto S., Yabuzaki J., Kanehisa M., Yamashita A., Oshima K., Furuya K., Yoshino C., Shiba T., Hattori M., Ogasawara N., Hayashi H., Hiramatsu K. comments: [('STRAIN', 'N315')] location: Lancet 357:1225-1240(2001).

positions: ['NUCLEOTIDE SEQUENCE [LARGE SCALE GENOMIC DNA].']

references: [('PubMed', '11418146'), ('DOI', '10.1016/S0140-6736(00)04403-2')]

title: Whole genome sequencing of meticillin-resistant Staphylococcus aureus.

Authors: Scherl A., Francois P., Bento M., Deshusses J.M., Charbonnier Y., Converset V., Huyghe A., Walter N., Hoogland

C., Appel R.D., Sanchez J.-C., Zimmermann-Ivol C.G., Corthals G.L., Hochstrasser D.F., Schrenzel J.

comments: [('STRAIN', 'N315')]

location: J. Microbiol. Methods 60:247-257(2005).

positions: ['IDENTIFICATION BY MASS SPECTROMETRY.']

references: [('PubMed', '15590099'), ('DOI', '10.1016/j.mimet.2004.09.017')]

title: Correlation of proteomic and transcriptomic profiles of Staphylococcus aureus during the post-exponential phase

of growth.

Authors: Vaezzadeh A.R., Deshusses J., Lescuyer P., Hochstrasser D.F.

comments: [('STRAIN', 'N315')]

location: Submitted (OCT-2007) to UniProtKB.

number: 3

positions: ['IDENTIFICATION BY MASS SPECTROMETRY [LARGE SCALE ANALYSIS].']

references: []

title: Shotgun proteomic analysis of total and membrane protein extracts of S. aureus strain N315.

seqinfo: (159, 18251, '811898409FEAFAAB'

MTLSILVAHDLQRVIGFENQLPWHLPNDLKHVKKLSTGHTLVMGRKTFESIGKPLPNRRNVVLTSDTSFNVEGVDVIHSIEDIYQLPGHVFIFGGQTLFEEMIDKVDDMYITVIEGKFRGDTFFPPYTFEDWEVASSVEGKLDEKNTIPHTFLHLIRKK

sequence_length: 159

sequence_update: ('23-JAN-2007', 2)

taxonomy_id : ['158879']

Glutamate racemase - UniProt

number: 1

accessions: ['P63638', 'Q99UV6'] annotation_update: ('05-DEC-2018', 85) comments: ['FUNCTION: Provides the (R)-glutamate required for cell wall biosynthesis. {ECO:0000255 | HAMAP-Rule:MF_00258}.', 'CATALYTIC ACTIVITY: Reaction=L-glutamate = D-glutamate; Xref=Rhea:RHEA:12813, ChEBI:CHEBI:29985, ChEBI:CHEBI:29986; EC=5.1.1.3; Evidence={ECO:0000255|HAMAP-Rule:MF_00258};', 'PATHWAY: Cell wall biogenesis; peptidoglycan biosynthesis. {ECO:0000255|HAMAP-Rule:MF_00258}.', 'SIMILARITY: Belongs to the aspartate/glutamate racemases family. {ECO:0000255|HAMAP-Rule:MF_00258}.'] created: ('11-OCT-2004', 0) cross_references: [('EMBL', 'BA000018', 'BAB42246.1', '-', 'Genomic_DNA'), ('PIR', 'B89886', 'B89886'), ('RefSeq', "WP_001039659.1', 'NC_002745.2'), ('ProteinModelPortal', 'P63638', '-'), ('SMR', 'P63638', '-'), ('EnsemblBacteria', 'BAB42246', 'BAB42246', 'BAB42246'), ('KEGG', 'sau:SA0997', '-'), ('HOGENOM', 'HOG000262396', '-'), ('KO', 'K01776', '-'), ('OMA', 'VYGCTHY', '-'), ('BioCyc', 'SAUR158879:G1G21-1138-MONOMER', '-'), ('UniPathway', 'UPA00219', '-'), ('Proteomes', 'UP000000751', 'Chromosome'), ('GO', 'GO:0008881', 'F:glutamate racemase activity', 'IEA:UniProtKB-UniRule'), ('GO', 'GO:0071555', 'P:cell wall organization', 'IEA:UniProtKB-KW'), ('GO', 'GO:0009252', 'P:peptidoglycan biosynthetic process', 'IEA:UniProtKB-UniRule'), ('GO', 'GO:0008360', 'P:regulation of cell shape', 'IEA:UniProtKB-KW'), ('HAMAP', 'MF_00258', 'Glu_racemase', '1'), ('InterPro', 'IPR015942', 'Asp/Glu/hydantoin_racemase'), ('InterPro', 'IPR001920', 'Asp/Glu_race'), ('InterPro', 'IPR018187', 'Asp/Glu_racemase_AS_1'), ('InterPro', 'IPR033134', 'Asp/Glu_racemase_AS_2'), ('InterPro', 'IPR004391', 'Glu_race'), ('PANTHER', 'PTHR21198:SF2', 'PTHR21198:SF2', '1'), ('Pfam', 'PF01177', 'Asp_Glu_race', '1'), ('SUPFAM', 'SSF53681', 'SSF53681', '2'), ('TIGRFAMs', 'TIGR00067', 'glut_race', '1'), ('PROSITE', 'PS00923', 'ASP_GLU_RACEMASE_1', '1'), ('PROSITE', 'PS00924', 'ASP_GLU_RACEMASE_2', '1')] data class: Reviewed description: RecName: Full=Glutamate racemase {ECO:0000255 | HAMAP-Rule:MF 00258}; EC=5.1.1.3 {ECO:0000255 | HAMAP-Rule:MF_00258}; entry_name: MURI_STAAN features : [('CHAIN', 1, 266, 'Glutamate racemase.', 'PRO_0000095509'), ('REGION', 9, 10, 'Substrate binding. {ECO:0000255|HAMAP-Rule:MF_00258}.', "), ('REGION', 41, 42, 'Substrate binding. {ECO:0000255|HAMAP-Rule:MF_00258}.', "), ('REGION', 73, 74, 'Substrate binding. {ECO:0000255 | HAMAP- Rule:MF_00258}.', "), ('REGION', 185, 186, 'Substrate binding. {ECO:0000255|HAMAP- Rule:MF 00258}.', "), ('ACT SITE', 72, 72, 'Proton donor/acceptor. {ECO:0000255|HAMAP-Rule:MF_00258}.', "), ('ACT_SITE', 184, 184, 'Proton donor/acceptor. {ECO:0000255|HAMAP-Rule:MF_00258}.', ")] gene_name: Name=murl {ECO:0000255|HAMAP-Rule:MF_00258}; OrderedLocusNames=SA0997; host organism:[] host taxonomy id:[] keywords: ['Cell shape', 'Cell wall biogenesis/degradation', 'Complete proteome', 'Isomerase', 'Peptidoglycan synthesis'] molecule_type: None organelle: organism: Staphylococcus aureus (strain N315). organism_classification: ['Bacteria', 'Firmicutes', 'Bacilli', 'Bacillales', 'Staphylococcaceae', 'Staphylococcus'] protein_existence: 1 references: Authors: Kuroda M., Ohta T., Uchiyama I., Baba T., Yuzawa H., Kobayashi I., Cui L., Oguchi A., Aoki K., Nagai Y., Lian J.-Q., Ito T., Kanamori M., Matsumaru H., Maruyama A., Murakami H., Hosoyama A., Mizutani-Ui Y., Takahashi N.K., Sawano T., Inoue R., Kaito C., Sekimizu K., Hirakawa H., Kuhara S., Goto S., Yabuzaki J., Kanehisa M., Yamashita A., Oshima K., Furuya K., Yoshino C., Shiba T., Hattori M., Ogasawara N., Hayashi H., Hiramatsu K. comments: [('STRAIN', 'N315')] location: Lancet 357:1225-1240(2001).

positions: ['NUCLEOTIDE SEQUENCE [LARGE SCALE GENOMIC DNA].']

references: [('PubMed', '11418146'), ('DOI', '10.1016/S0140-6736(00)04403-2')]

title: Whole genome sequencing of meticillin-resistant Staphylococcus aureus.

Authors: Vaezzadeh A.R., Deshusses J., Lescuyer P., Hochstrasser D.F.

comments: [('STRAIN', 'N315')]

location: Submitted (OCT-2007) to UniProtKB.

number: 2

positions: ['IDENTIFICATION BY MASS SPECTROMETRY [LARGE SCALE ANALYSIS].']

references: []

title: Shotgun proteomic analysis of total and membrane protein extracts of S. aureus strain N315.

seqinfo: (266, 29698, '93BC35DB4A2C7B0E')

sequence:

MNKPIGVIDSGVGGLTVAKEIMRQLPNETIYYLGDIGRCPYGPRPGEQVKQYTVEIARKLMEFDIKMLVIACNTATAVALEYLQKTLSIPVIGVIEPGART AIMTTRNQNVLVLGTEGTIKSEAYRTHIKRINPHVEVHGVACPGFVPLVEQMRYSDPTITSIVIHQTLKRWRNSESDTVILGCTHYPLLYKPIYDYFGGKKT VISSGLETAREVSALLTFSNEHASYTEHPDHRFFATGDPTHITNIIKEWLNLSVNVERISVND

sequence_length: 266

sequence_update : ('11-OCT-2004', 1)

taxonomy_id: ['158879']