The Protein Data Bank: A Computer-based Archival File for Macromolecular Structures

The Protein Data Bank is a computer-based archival file for macromolecular structures. The Bank stores in a uniform format atomic co-ordinates and partial bond connectivities, as derived from crystallographic studies. Text included in each data entry gives pertinent information for the structure at hand (e.g. species from which the molecule has been obtained, resolution of diffraction data, literature citations and specifications of secondary structure). In addition to atomic co-ordinates and connectivities, the Protein Data Bank stores structure factors and phases, although these latter date are not placed in any uniform format. Input of data to the Bank and general maintenance functions are carried out at Brookhaven National Laboratory. All data stored in the Bank are available on magnetic tape for public distribution, from Brookhaven (to laboratories in the Americas), Tokyo (Japan), and Cambridge (Europe and worldwide). A master file is maintained at Brookhaven and duplicate copies are stored in Cambridge and Tokyo. In the future, it is hoped to expand the scope of the Protein Data Bank to make available co-ordinates for standard structural types (e.g. α-helix, RNA double-stranded helix) and representative computer programs of utility in the study and interpretation of macromolecular structures.

The Protein Data Bank† (1971,1973) was established in 1971 as a computer-based archival file for macromolecular structures. The purpose of the Bank is to collect, standardize, and distribute atomic co-ordinates and other data from crystallographic studies. As the number of solved protein and nucleic acid structures has grown to the point where some 10° characters are necessary to represent the co-ordinate information currently held, the need for such a computer-readable file has become very clear, and demands for the Bank's services have increased accordingly. The Protein Data Bank is one of several data base activities in the field of crystallography, e.g. the Bibliographic (Kennard et al., 1972) and Structural (Allen et al., 1973) Data Files for organic and organometallic compounds, the Atlas of Macromolecular Structure on Microfiche (AMSOM) (Feldmann, 1977), the Bond Index to the Determination of Inorganic Crystal Structures (BIDICS)‡ and the Powder Diffraction File.§

(a) Scope

The Protein Data Bank covers atomic co-ordinates, structure factors and phases from diffraction studies of macromolecules. Since most of this information is not generally published in the primary literature, the Bank depends for comprehensiveness on data supplied directly by the investigators. It is essentially a depository of data, held in computer-readable form, in contrast to other data banks that are based

[§] American Society for Testing Materials, 1916 Race St., Philadelphia, PA. 19103, U.S A.



[†] Protein Data Bank is a misnomer of historical origin, since the file now contains entries for a nucleic acid.

[‡] I. D. Brown, Bond Index to the Determination of Inorganic Crystal Structures, McMaster University, Hamilton, Ontario, Canada, L8S 4M1.

TABLE 1

Protein data bank holdings

| | | 4 | |
|-----------------|--|---|----------|
| IDENT | MOLECULE ADENYLATE KINASE ALCONOL DENYDROGENASE (ADP-RIB) ALCONOL DENYDROGENASE (ORTHOPNEN) BENCE-JONES INTUMOGLOBUL IN REI CALCIUM-BINDING FRAGRIENT (NEW) BENCE-JONES INTUMOGLOBUL IN REI CALCIUM-BINDING PARVALBUMIN SET 6A CALCIUM-BINDING PARVALBUMIN SET 6A CALCIUM-BINDING PARVALBUMIN SET 6A CARCONIC ANNYDRASE B CARRONIC ANNYDRASE B CARRONIC ANNYDRASE C CARBOXYPEPTIDASE A CONCANAVELIN A CONCANAVE CONCANAVE CONCANA CO | DEPOS & TOR | STATUS |
| IADK | ADENY ATE KINASE | G. SCHULZ | A |
| 2ADH | ALCOHOL DENYDROGENASE (ORTHOPHEN) | CI. BRANDEN | |
| 3CHA | ALPHA-CHYMOTRYPS IN | A. TUL INSKY | K |
| IREI | BENCE-JONES INTUNOGLOBULIN REI | O. EPP. R. HUBER | |
| 2CPV | CALCIUM-BINDING PARVALBUMIN SET 6A CALCIUM-BINDING PARVALBUMIN SET 6N | R. KRETSINGER R. KRETSINGER | |
| 3CPV 1CAB | CALCIUM-BINDING PARVALBUMIN SET 61 CARBONIC ANNYDRASE B | R. KRETSINGER K. KANNAN | |
| ICAC | CARBONIC ANNYDRASE C CARBOXYPEPTIDASE A | K. KANNAN U. LIPSCOMB | |
| LCHG | CHYMOTRYPS INOGEN | J. KRAUT G. REEKE. G. EDELMAN | N |
| 3CHA IBSC | CONCANAVALIN A | K. HARDNAH F. S. MATHEUS | R |
| ICYT | CYTOCHRONE C (ALBACORE, OXIDIZED) | R. DICKERSON | |
| วัยวัติ เกรา | CYTOCHRONE C (BONITO, HEART) | M. KAKUDO | |
| 155C | CYTOCHRONE C550 | R. TIMKOVICH | |
| IFDX | FERREDOXIN | L. JEHSEH | |
| IGCH | GATTA-CHYPETRYPSIN | COHEN. DAVIES. SILVERTON | P |
| 2HHB | HEIMGLOBIH (HORSE, AQUO MET) | LADNER, HEIDHER, PERUTA | Z RP |
| IHHB | HENOGLOBIH (HUNAH, DEOXY) | M. PERUTZ. G. FERMI | |
| ILHB | HENDGLOBIN (HONAN' LEINT' DEOXA) | W. HENDRICKSON | |
| INIP | HEXUKINASE (YEAST) BIII HIGH POTENTIAL IRON PROTEIN | J. KRAUT | В |
| 2LDH 31.DH | LACTATE DENYDROGENASE LACTATE DENYDROGENASE/MAD/PYRUVATE | M. ROSSMANN M. ROSSMANN | PD PD |
| ILYZ 2LYZ | LYSOZYME (HEN EGG-WHITE, SET W2) LYSOZYME (HEN EGG-WHITE, SET RSSD) | R. DIAMOND R. DIAMOND | P P: |
| 3ĽÝŽ 4ĽÝŽ | LYSOZYME (NEN EGG-WHITE, SET RSGA) LYSOZYME (NEN EGG-WHITE, SET RS9A) | R. DIAMOND | P |
| SLYZ 61 YZ | LYSOZYNE (NEN EGG-UHITE, SET RS12A) | R. DIAMOND R. DIAMOND | P |
| HOMI Heat | MALATE DENYDROGENASE HYOGI OBIN (SPERM LING) F) | L. BANASZAK | Ä |
| 21BN 31BN | NYOGLOBIN (SPERM WHALE, MET) | T. TAKAHO | |
| 3PT I | PANCREGIEC TRIPSIN INHIBITOR | R. HUBER | R |
| 2PAP | PAPAIN (ACE-ALA-ALA-PHE-ALA, CYS-25) | J. DRENTH | K |
| 4PAP | PAPAIN (OXIDIZED CYS-25) | J. DRENTH | |
| 6PAP | PAPAIN (BZOXY-GL Y-PNE-GLY, CYS-25) | J. DRENTH | |
| IPGK | PHOSPHOGLYCERATE KINASE (YEAST) | H. WATSON | A |
| IPAB | PREALBUMIN (NUMAN, PLASMA) | S. OATLEY, D. PHILLIPS | |
| IRHS 2RXM | RIBONUCLEASE S RUBREDOXIN | H. LOYCKOFF L. JENSEN | *IB |
| 1SHS 1SGB | STAPHYLOCOCCAL NUCLEASE STREPTOMYCES GRISEUS PROTEINASE B | F. A. COTTON, E. HAZEN M. JAMES | А |
| ISBT 2SBT | SUBTILISIN BPN' SUBTILISIN HOVO | J. KRAUT J. DRENTH | |
| ISOD | SUPEROXIDE DISMUTASE THERMOLYSIN (UNREFINED) | J. AND D. RICHARDSON | A |
| 211.H !SRX | THERMOLYSIN (REFINED) | B. MATTHELIS BO. SODERBERG | А |
| ITHA 21HA | TRANSFER RNA (YEAST, PHE) | J. SUSSMAN, SH. KIM M. SUNDARALINGAM | H |
| 31HA | TRANSFER RNA (YEAST, PNE) TRANSFER RNA (YEAST, PNE) TRANSFER RNA (YEAST, PNE) | JACK, LADHER, KLUG | P |
| ITIM | TRIOSE PHOSPHATE ISOMERASE TRYPSIN (HATIVE, PHB) | FEHLHAMMER.BODE.SCHUNG | ER H |
| 2PTB IPTC | TRYPSIN(BEHZAMIDINE INNIBITED, PH?) TRYPSIN/TRYPSIN INNIBITOR COMPLEX | FENLHAMMER, BODE, SCHWAG BODE ET AL. | ER RH |
| STA | IVS CODES | | |
| BLANK | STANDARD ENTRY AVAILABLE FOR DISTRIBU | UTION | |
| D D | ALPHA CARBON ATOR'S OHLY BACKBONE ONLY HEW DATA HAS BEEN PROMISED | | |
| ň P | NEW ENTRY WITH DEPOSITOR FOR APPROVAL IN PREPARATION | L | |
| R | REPLACES AN OUT OF DATE PARAMETER SE | T | |



on data abstracted from scientific publications. The Bank contains 77 atomic coordinate entries for 47 macromolecules (Table 1),† and 13 sets of structure factors and phases. The atomic co-ordinate entries, which include descriptive text and partial bond connectivities, conform to a uniform format (see below), but the structure factors and phases are stored in the format received from depositors. All co-ordinate entries are referred to depositors for verification, before being made available publicly through the Bank.

(b) Record structure of atomic co-ordinate entries

Atomic co-ordinate entries consist of records each of 80 characters.‡ Using the punched card analogy, columns 1 to 6 contain a record type identifier, and columns 7 to 70 contain data.§ Columns 71 to 80 are normally blank, but may contain sequence information which is added by the library-file management program UPDATE¶ used to maintain the file on the Brookhaven CDC CYBER 70/76 computing system. In order to facilitate retrieval of data from the file, the first four characters of each record define the unique record type, and the syntax of each record is independent of the order of records within any entry for a particular macromolecule. (In the master file, this order is always fixed.) Atomic co-ordinate data contributed by depositors are processed into the standard format with program MACMOL, || which also subjects the data to certain nomenclature and connectivity checking procedures.

A sample partial entry for the protein ribonuclease S is shown in Table 2.†† The unique code 1RNS identifying this entry is given in the HEADER record, along with the date these data were entered into the Bank, and a provisional classification based on function, intended for future use in indexing and subdividing the file. Text giving the name of molecule, species from which it has been obtained, authors, literature citations, and other general description are presented in records COMPND through REMARK. SEQRES gives the amino acid sequence, and FTNOTE records are footnotes keyed to particular residues or atoms. Records HELIX through TURN describe the secondary structure as stated or approved by the depositor. Record CRYST1 defines the unit cell, while ORIGX and SCALE respectively give transformations relating the orthogonal Angström co-ordinates stored in the file to those originally supplied by the depositor (these frequently are referred to an oblique or non-isometric system) and to standard crystallographic fractional co-ordinates. ATOM records give the IUPAC-IUB (1969) standard atom names (IUPAC-IUB, 1970), and residue abbreviations (IUPAC-IUB, 1971), along with sequence identifiers (cf. SEQRES, above), co-ordinates in Angström units, and occupancies and thermal

- † In addition to current co-ordinate entries shown in Table 1, the Bank contains obsolete entries (for adenylate kinase tosyl, α -chymotrypsin, concanavalin A, lactate dehydrogenase, horse methemoglobin, papain, rubredoxin, benzamidine-inhibited trypsin and pancreatic trypsin inhibitor), which have been superseded by later, more accurate data. These obsolete data are available on special request.
- † Originally, the Bank used a 140-character format, similar to that employed in the protein refinement programs of Diamond (1966,1971). The 140-character format has been superseded by the 80-character format.
 - § A detailed description of the file formats is available from Brookhaven on request.
- ¶ Control Data Corporation, UPDATE Reference Manual, Publication No. 60342500, Control Data Corporation, Arden Hills, Minnesota, 1974.
- G. J. B. Williams, unpublished. For the 140-character data, program PROIN by E. F. Meyer was utilized.
- †† The file is organized in a similar way for proteins and nucleicacids, althoughcertain differences exist, e.g. with regard to details of atom and residue names.



Table 2
Abbreviated sample atomic co-ordinate entry (ribonuclease S)

```
HEADER
            HYOROLASE (PHOSPHORIC DIESTER, RNA)
                                                          01-APR-73
                                                                        IRNS
            RIBONUCLFASE-S (E.C. 3.1.4.22)
BOVINE (ROS TAURUS) PANCREAS
COMPNO
SOURCE
            F. M. RICHAROS AND H. W. WYCKOFF PRELIMINARY REFINEMENT
AUTHOR
JRNL
            OF PROTETH COORDINATES IN REAL SPACE. ACTA CRYST. . VOL. A31.
JRNL
JRNL
            P698 (1975).
REMARK
          1 REFERENCE 1. F. M. RICHARDS AND H. W. WYCKOFF, ATLAS OF STRUCTURES FOR MOLECULAR BIOLOGY, VOL. 1. RIBONUCLEASE-S.
REMARK
REMARX
REMARK
              CLARENDON PRESS (1973).
          1 REFERENCE 2. F. M. RICHARDS AND H. W. WYCKOFF, BOVINE
REMARK
          1 PANCREATTC RIBONUCLEASE, THE ENZYMES, EDITEU BY P. O.
REMARK
          1 BOYER FOL. IV. THIRD EDITION. P647. ACADEMIC PRESS (1971)
1 REFERENCE 3. F. M. RICHARDS, H. W. WYCKOFF. W. D. CARLSON.
REMARK
REMARK
             N. M. ALLEWELL, B. LEE AND Y- MITSUI, PROTEIN STRUCTURE, RIBONUCLEASE-S AND NUCLEOTIDE INTERACTIONS, COLO SPRING
REMARK
REMARK
REMARK
              HARBOR SYMPOSIA ON QUANTITATIVE BIOLOGY. VOL. XXXVI. P35
REMARK
REMARK
          1 REFERENCE 4. N. M. ALLEWELL AND H. W. WYCKOFF.
             CRYSTALI OGRAPHIC ANALYSIS OF THE INTERACTION OF CUPRIC ION WITH RIBONUCLEASE S. J. BIOL. CHEM. VOL. 246. P4657
REMARK
REMARK
REMARK
              (1971)
REMARK
          1 REFERENCE S. H. W. WYCKOFF, D. TSEHNOGLOU, A. W. HANSON,
              J. R. KNOX. B. LEE AND F. M. RICHARDS. THE THREE-
DIMENSTANAL STRUCTURE OF RIBDNUCLEASE-S. INTERPRETATION
REMARK
REMARK
              OF AN FLECTRON DENSITY MAP AT A NOMINAL RESOLUTION OF 2
REMARK
REMARK
              ANGSTROMS. J. BIOL. CHEM. . VOL. 245. P305 (1970).
REMARK
          1 REFERENCE 6. H. W. WYCKOFF, K. D. HAROMAN, N. M. ALLEWELL,
REMARK
              T. INAGAMI. D. TSERNOGLOU. L. N. JOHNSON AND F. M.
             RICHARDS. THE STRUCTURE OF RIBONUCLEASE-S AT 6 ANGSTROM
REMARK
REMARK
             RESOLUTION, J. BIOL. CHEM. . VOL. 242. P3749 (1967) .
REMARK
REMARK
          2 RESOLUTION. 2.0 ANGSTROMS.
REMARK
REMARK
          3 REFINEMENT. BY A STEEPEST-DESCENTS PROCEDURE. REFER TO THE
REMARK
            JRNL CITATION ABOVE.
REMARK
REMARK
          4 THIS COMPDINATE SET IS DESIGNATED BY THE DEPOSITOR.
REMARK
REMARK
          5 THE *S-PEPTIDE* (RESIOUES 1-20) WHICH FORMS A SEPARATE
          5 CHAIN FROM THE REMAINDER OF THE MOLECULE IS GIVEN THE
REMARK
REMARK
          5 CHAIN INFETIFIER S.
SEGRES
          1 5
                20 LYS GLU THR
                                   ALA ALA ALA LYS PHE GLU ARG GEN HIS MET
SEGRES
          2 5
                 20
                     ASP SER SER THR SER ALA ALA
SEGRES
                104
                      FER SER SER ASN TYR CYS ASN GLN MET MET LYS SER ARG
SEGRES
                104
                      ASN LEU THR LYS ASP ARG CYS LYS PRO VAL ASN THR PHE
SEGRES
                104
                     VAL HIS GLU SER LEU ALA ASP VAL GLN ALA VAL CYS SER
SEGRES
                104
                     RLY LYS ASN VAL ALA CYS LYS ASN GLY GLN THR ASN CYS
SEGRES
                104
                     TYR GLN SER TYR SER THR MET SER ILE THR ASP CYS ARG
SEGRES
                104
                     ALU THR GLY SER SER LYS TYR PRO ASH CYS ALA TYR LYS
SEORES
                104
                     THR THR GLN ALA ASN LYS HIS THE THE VAL ALA CYS GLU
SEGRES
                104
                     GLY ASN PRO TYR VAL PRO VAL HIS PHE ASP ALA SER VAL
FTNOTE
FTNOTE
          1 THE MAIN CHAIN AND HOST OF THE ASSOCIATED SIDE CHAINS ARE
FTNOTE
          1 NOT WELL-DEFINED IN THE REGIONS OF RESIDUES 2, 65-72 AND
FTNOTE
          1 119-123.
FTNOTE
FTNOTE
            THE MAIN CHAIN IS VERY POORLY DEFINED OR NOT VISIBLE AT ALL
FTNOTE
          2 IN THE FLECTRON DENSITY MAP IN THE REGIONS OF RESIDUES 1. 2 18-20.21-23 AND 124. 1 H1 THR 5 3 MET S 13 1
FTNOTE
             H1 THR 5
HELIX
HELIX
             HZ ASN
                          24 ASN
                                        34
```



TABLE 2-continued

| HELIX | 3 | H3 S | | 50 | ALA | 56 | | | | | | | | | |
|---------------------------|-----|---------|------|------|------|---------|------|------|------|--------|------|-----|--------|----|----|
| SHEET | 1 | | LYS | 41 | HIS | | _ | | | | | _ | | | _ |
| SHEET | 2 | | MFT | 79 | THE | | -1 | | AS | | 44 | 0 | CYS | | 84 |
| SHEET | 3 | 51 3 | ALA | 96 | LYS | 104 | -1 | N | AS | P | 43 | 0 | THR | 1 | 00 |
| SHEET | 1 | 52 4 | | 61 | ALA | | | | | | | | | | |
| SHEET | 2 | 52 4 | | T1 | SER | 75 | -1 | N | VA | | 63 | 0 | CYS | | 72 |
| SHEET | 3 | 52 4 | HIS | 105 | GLU | 111 | -1 | | TY | | 73 | 0 | VAL | 1 | 00 |
| SHEET | 4 | 52 4 | VAL | 116 | VAL | 124 | -1 | Q | AL | . A | 109 | N | VAL | 1 | 18 |
| TURN | 1 | TI V | AL | | VAL | 57 | | PSEL | DO | 3/10 | HEL | KI | | | |
| TURN | 2 | TZ A | _ | | SER | 59 | | PSEL | 100 | 3/10 | HEL | IX | | | |
| TURN | 3 | T3 C | | | GL Y | 68 | | BETH | 1 51 | RNDS | 1.2 | OF | SHEET | 52 | |
| TURN | 4 | T4 T | | | SER | 90 | | ENO | OF | STRA | NO 2 | OF | SHEET | 51 | |
| CRYSTI | | .650 | | 650 | | | .04 | | | | | | 2 1 | | 6 |
| ORIGXI | 77 | | 0000 | 0.00 | | 0.000 | | | | | 0000 | | | | |
| ORIGX2 | | | 0000 | 1.00 | | 0.000 | | | | | 0000 | | | | |
| ORIGXE | | | 000n | | 0000 | 1.000 | | | | U - OC | 0000 | | | | |
| SCALEI | | | 2306 | | 2931 | 0.000 | | | | | 0000 | | | | |
| | | | 0000 | | 5861 | 0.000 | | | | | 0000 | | | | |
| SCALE2 | | | 0000 | 0.00 | | .010 | | | | | 0000 | | | | |
| SCALES | 1 | | | | 0000 | -15.39 | | 7.9 | 416 | | 202 | 1.0 | 0 0- | 00 | 2 |
| HOTA | 1 | N CA | LYS | | | -15.14 | | 7.6 | | | 730 | 1.0 | | 00 | 2 |
| HOTA | 3 | CA | LYS | - | | -14.96 | | | 107 | | 763 | 1.0 | | 00 | 2 |
| ATOH | 4 | 0 | LYS | | | -15.14 | | 5.3 | | | 732 | 1.0 | | 00 | 2 |
| ATOH | 5 | CB | LYS | | | -13.87 | | | 244 | | 185 | 1.0 | | 00 | 2 |
| ATOH | 6 | CG | LYS | | | -12.69 | | 7.6 | | | 794 | 1.0 | | 00 | 2 |
| ニョン コ レきれかしきんきん | _ | | | - | | | | | | | | | せいめいせい | | |
| ATOM | 927 | N N | ASP | 121 | | -6.79 | 15 | -9.2 | | 7. | 034 | 1.0 | | 00 | 1 |
| HOTA | 928 | CA | ASP | 121 | | -5.81 | | -9.4 | | | 935 | 1.0 | | 00 | ī |
| ATOM | 929 | C | ASP | 121 | | -6.21 | 7 - | 10.1 | 156 | | 789 | 1.0 | 0 0. | 00 | 1 |
| POTA | 930 | 0 | ASP | 121 | | -5.82 | | -9.6 | 150 | | 652 | 1.0 | 0 0. | 00 | 1 |
| POTA | 931 | CB | ASP | 121 | | -4.52 | | | | | 648 | 1.0 | | 00 | 1 |
| ATOH | 932 | CG | ASP | 121 | | -3.47 | 1 | -4.5 | 503 | 5. | 687 | 1.0 | 0 0. | 00 | 1 |
| ATOH | 933 | | ACD | 121 | | -3.32 | | -8.0 | | | 636 | 1.0 | | 00 | ī |
| ATOM | 934 | | ASP | 121 | | -2.7 | | | | | 799 | 1.0 | | 00 | 1 |
| ATOH | 935 | N | ALA | 122 | | -7.04 | | | | | 013 | 1.0 | | 00 | 1 |
| HOTA | 936 | CA | AL A | 122 | | -7.96 | 5 - | 12.0 | 186 | 4. | 084 | 1.0 | 0 0. | 00 | 1 |
| ATOH | 937 | C | ALA | 122 | | -8.55 | | | | | 724 | 1.0 | | 00 | 1 |
| ATOH | 938 | 0 | AL A | 122 | | -8.49 | 95 - | 13.6 | 536 | | 925 | 1.0 | 0 0. | 00 | 1 |
| HOTA | 939 | CB | A A | 122 | | -6.99 | 1 - | 12.5 | 510 | 2. | 881 | 1.0 | 0 0. | 00 | 1 |
| ATOM | 940 | N | SER | 123 | | -0.00 | | | | | 717 | 1.0 | 0 0. | 00 | 1 |
| ATOH | 941 | CA | SFP | 123 | | -9.75 | 68 - | 15.1 | 155 | 3. | 627 | 1.0 | 0 0. | 00 | 1 |
| MOTA | 942 | C | SFp | 123 | | -8.91 | 5 - | 16.1 | 27 | | 880 | 1.0 | 0 0. | 00 | 1 |
| HOTA | 943 | 0 | SFR | 123 | | -8.37 | 2 - | 15.8 | 312 | 1. | 810 | 1.0 | | 00 | 1 |
| ATOM | 944 | CB | SFR | 123 | | -10.87 | | | | | 597 | 1.0 | | 00 | 1 |
| HOTA | 945 | OG | SER | 123 | | -10.19 | | | | | 530 | 1.0 | | 00 | 1 |
| HOTA | 946 | N | VAL | 124 | | -8 . B4 | | | | | 438 | 1.0 | | 00 | 2 |
| HOTA | 947 | CA | VAL | 124 | | -8.59 | | | | | 596 | 1.0 | | 00 | 2 |
| ATOM | 948 | C | VAL | 124 | | -9.23 | | | | | 209 | 1.0 | 0 0- | 00 | 2 |
| ATOH | 949 | 0 | VAL. | 124 | | -8.56 | | | | | 377 | 1.0 | | 00 | 2 |
| ATOM | 950 | CB | VAL | 124 | | -8.93 | | | | | 162 | 1.0 | | 00 | 2 |
| HOTA | 951 | CGI | | 124 | | -9.13 | | | | | 012 | 1.0 | | 00 | 2 |
| POTA | 952 | | VAL | 124 | | -7.76 | | | | | 226 | 1.0 | | 00 | 2 |
| ATOH | 953 | OXT | | 124 | | -10.41 | | | | | 046 | 1.0 | | 00 | 2 |
| TER | 954 | | VAL. | 124 | | | | | - | | | | | | _ |
| CONECT | 196 | 195 | 644 | | | | | | | | | | | | |
| CONECT | 312 | 311 | | | | | | | | | | | | | |
| CONECT | 448 | 447 | | | | | | | | | | | | | |
| CONECT | 499 | 497 | | | | | | | | | | | | | |
| CONECT | 549 | 498 | | | | | | | | | | | | | |
| CONECT | 644 | 196 | | | | | | | | | | | | | |
| CONECT | 729 | 312 | | | | | | | | | | | | | |
| CONECT | 844 | 448 | | | | | | | | | | | | | |
| | - | | | 0 | 3 | 7 | | | 0 | 6 | 952 | | 2 | 8 | 10 |
| MASTER | | 36 | 1 0 | - | | | | | | | | | | | |



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