# Membrane and Soluble Proteins 

The Morpheus® Additive Screen MD1-93
The Morpheus Additive Screen is a new 96-condition additive screen, containing an extensive library of ligands suitable for soluble and membrane proteins.

## MD1-93 is presented as $96 \times 100 \mu \mathrm{~L}$ conditions.

Features of The Morpheus Additive Screen:

- Test a large range of precipitants, stabilizers, buffers, nucleants, phasing compounds, cryoprotectants and surfactants all in one screen.
- Enhance stability and solubility of protein for crystallization with the inclusion of NDSBs, polyamines, monosaccharides and amino acids.
- Contains heavy atoms as additives for experimental phasing.
- Use in protein stability screen assays e.g. thermal shift assays.


## Introduction

The Morpheus Additive Screen contains a library of additives for use in protein crystallization optimization experiments (Table $1 \& 2)$.
These additives are a mixture of low molecular weight components typically found in many solved protein structures and have been found to aid with crystallization. A range of less typical additives which are meant to alter protein stability and solubility such as heavy metals, NDSBs, polyamines, monosaccharides and amino acids are also included.

[^0]
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## Formulation Notes:

The Morpheus Additive Screen reagents are formulated using ultrapure water ( $>18.0 \mathrm{M} \Omega$ ) and are sterilefiltered using $0.22 \mu \mathrm{~m}$ filters. No preservatives are added. Material Safety Datasheets are available from our website.

Final pH may vary from that specified on the datasheet. Molecular Dimensions will be happy to discuss the precise formulation of individual reagents.

Individual reagents and stock solutions for optimization are available from Molecular Dimensions.
Enquiries regarding The Morpheus Additive Screen formulation, interpretation of results or optimization strategies are welcome. Please e-mail, fax or phone your query to Molecular Dimensions.
Contact and product details can be found at www.moleculardimensions.com

RE-ORDERING INFORMATION

| Code | Pack Size | Description |
| :---: | :---: | :---: |
| MD1-46 | $96 \times 10 \mathrm{~mL}$ | Morpheus |
| MD1-47 | $96 \times 1 \mathrm{~mL}$ | Morpheus HT-96 |
| MD1-47-FX | 96x $100 \mu \mathrm{~L}$ | Morpheus FX-96 pre-filled plate |
| Other Morpheus screens |  |  |
| MD1-91 | $96 \times 10 \mathrm{~mL}$ | Morpheus II |
| MD1-92 | $96 \times 1 \mathrm{~mL}$ | Morpheus II HT-96 |
| MD1-92-FX | 96x $100 \mu \mathrm{~L}$ | Morpheus II FX-96 pre-filled plate |
| MD1-93 | $48 \times 100 \mu \mathrm{~L}$ | The Morpheus® Additive screen |
| MD1-116 | $96 \times 10 \mathrm{~mL}$ | Morpheus III |
| MD1-117 | $96 \times 1 \mathrm{~mL}$ | Morpheus III HT-96 |
| MD1-118 | $48 \times 100 \mu \mathrm{~L}$ | Hippocrates ${ }^{\text {TM }}$ additive screen |
| Green screens (contain green fluorescent dye - ideal for UV) |  |  |
| MD1-46-GREEN | $96 \times 10 \mathrm{~mL}$ | Morpheus Green screen |
| MD1-47-GREEN | $96 \times 1 \mathrm{~mL}$ | Morpheus HT-96 Green screen |
| Combo Packs |  |  |
| MD1-76 | $192 \times 10 \mathrm{~mL}$ | Power combo value pack <br> (Morpheus + MIDASplus) |
| MD1-76-HT | $192 \times 1 \mathrm{~mL}$ | Power combo value pack HT-96 <br> (Morpheus + MIDASplus HT-96) |
| Single reagents |  |  |
| MDSR-46-tube number | 100 mL | Morpheus single reagents |
| MDSR-47-well number | 100 mL | Morpheus HT-96 single reagents |
| MDSR-91-tube number | 100 mL | Morpheus II single reagents |
| MDSR-92-well number | 100 mL | Morpheus II HT-96 single reagents |

For Morpheus mixes and stock solutions please visit the Optimization section on our website.

## ||||||||||||||||||||||||||

Morpheus, Morpheus II and Morpheus III have been designed and developed by Fabrice GORREC, in collaboration with the scientists at the Medical Research Council Laboratory of Molecular Biology (LMB) at Cambridge and is manufactured exclusively under license from LifeARC by Molecular Dimensions Limited.

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## Membrane and Soluble Proteins

Table 1: List of PDB ligands in Morpheus

| PDB Ligand name(s) | Class | PDB ID(s) | Number of Structures* |
| :---: | :---: | :---: | :---: |
| 1,2-Ethanediol (ethylene glycol) | Precipitant | EDO, EGL | 1081 |
| 1,2-Propanediol (enantiomers R and S) | Alcohols | PGO, PGR | 41 |
| 1,3-Propanediol | Alcohols | PDO | 7 |
| 1,4-Butanediol | Alcohols | BU1 | 11 |
| 1,6-Hexanediol | Alcohols | HEZ | 19 |
| 1-Butanol | Alcohols | 1BO | 7 |
| 2-(N-Morpholino)-ethane sulfonic acid (MES) | Buffer | MES | 315 |
| 2-Amino-2-hydroxymethyl-propane-1,3-diol (Tris) | Buffer | TRS | 334 |
| 2-Methyl-2,4-pentanediol (MPD, enantiomers $R$ and S) | Precipitant | MPD, MRD | 504 |
| 3-Morpholinopropane-1-sulfonic acid (MOPS) | Buffer | MPO | 21 |
| 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) | Buffer | EPE | 201 |
| Acetic acid, acetate, acetyl | Carboxylic acids | ACY, ACT, ACE | 1890 |
| (S)-2-Aminopropanoic acid (Alanine, (enantiomers L and D) | Amino acids | ALA, DAL | 35 |
| Amino, Ammonia, Ammonium | multiple | NH2, NH3, NH4 | 582 |
| N,N-bis(2-hydroxyethyl)glycine (Bicine) | Buffer | BCN | 13 |
| Bromide | Halogens | BR | 120 |
| Calcium | Divalents | CA | 3959 |
| Chloride | Multiple | CL | 2842 |
| Citric acid, citrate | Carboxylic acids | CIT, FLC | 384 |
| D-Galactose (anomers $\alpha$ and $\beta$ ) | Monosaccharides | GAL, GLA | 86 |
| D-Glucose (anomers $\alpha$ and $\beta$ ) | Monosaccharides | GLC, BGC | 206 |
| Glutamic acid (enantiomers L and D) | Precipitant | GLU, DGL | 75 |
| Di(Hydroxyethyl)ether (Di-Ethyleneglycol) | Ethylene glycols | PEG | 209 |
| D-Mannose (anomers $\alpha$ and $\beta$ ) | Monosaccharides | MAN, BMA | 178 |
| D-Xylopyranose (anomers $\alpha$ and $\beta$ ) | Monosaccharides | XYL, XYP | 41 |
| Fluoride | Halogens | F | 16 |
| Formic acid | Carboxylic acids | FMT | 267 |
| Glycerol | Amino acids | GOL | 2884 |
| Glycine | Buffer | GLY | 50 |
| Imidazole | Halogens | IMD | 154 |
| Iodide | Alcohols | IOD | 178 |
| Isopropyl alcohol (iso-propanol, 2-Propanol) | Monosaccharides | IPA, IOH | 174 |
| L-Fucose (anomers $\alpha$ and $\beta$ ) | Amino acids | FUC, FUL | 62 |
| Lysine (enantiomers L and D) | Amino acids | LYS, DLY | 36 |
| Magnesium | Divalents | MG | 3991 |
| N -Acetyl-d-glucosamine (anomers $\alpha$ and $\beta$ ) | Monosaccharides | NAG,NBG | 1150 |
| Nitrate | NPS | NO3 | 156 |
| Oxamic acid | Carboxylic acids | OXM | 17 |
| Penta(hydroxyethyl)ether (Penta-Ethyleneglycol) | Ethylene glycols | 1PE | 91 |
| Phosphates | NPS | PO4, PI, 2HP | 1687 |
| Potassium | Carboxylic acids | K | 720 |
| Serine (enantiomers L and D) | Amino acids | SER, DSN | 38 |
| Sodium | multiple | NA | 1926 |
| Sulfate | NPS | SO4 | 5793 |
| Tartaric acid (enantiomers R and S ) | Carboxylic acids | TAR, TLA | 113 |
| Tetra(hydroxyethyl)ether (Tetra-Ethyleneglycol) | Ethylene glycols | PG4 | 194 |
| Tri(Hydroxyethyl)ether (Tri-Ethyleneglycol) | Ethylene glycols | PGE | 107 |
|  32956 |  |  |  |

Table 2: List of PDB ligands in Morpheus II

| PDBligano nome | Closs | PDBID <br> (main) | No. of structures* |
| :---: | :---: | :---: | :---: |
| Lithium sulfate | Common salt | LI | 51 |
| Sodium chloride | Common salt | NA | 4726 |
| Potassium sulfate | Common salt | K | 1638 |
| Manganese chloride tetrahydrate | Divalent cation | MN | 1938 |
| Cobalt chloride hexahydrate | Divalent cation | CO | 474 |
| Nickel chloride hexahydrate | Divalent cation | NI | 699 |
| Zinc acetate dihydrate | Divalent cation | ZN | 8413 |
| Barium acetate | Alkali | BA | 91 |
| Cesium acetate | Alkali | CS | 75 |
| Rubidium chloride | Alkali | RB | 34 |
| Strontium acetate | Alkali | SR | 101 |
| Sodium chromate tetrahydrate | Oxometalate | CR | 7 |
| Sodium molybdate dihydrate | Oxometalate | MOO | 20 |
| Sodium orthovanadate | Oxometalate | VO4 | 73 |
| Sodium tungstate dihydrate | Oxometalate | WO4 | 47 |
| Erbium (III) chloride hexahydrate | Lanthanide | ER3 | 2 |
| Terbium (III) chloride hexahydrate | Lanthanide | TB | 11 |
| Ytterbium (III) chloride hexahydrate | Lanthanide | YB | 57 |
| Yttrium (III) chloride hexahydrate | Lanthanide | YT3 | 33 |
| Xylitol | Monosaccharide | XYL | 25 |
| D-(-)-fructose | Monosaccharide | FRU; FUD | 36; 4 |
| D-sorbitol | Monosaccharide | SOR | 12 |
| Myo-inositol | Monosaccharide | INS | 16 |
| L-rhamnose monohydrate | Monosaccharide | RAM | 43 |
| DL-threonine | Amino-acid | DTH; THR | 23; 1 /a |
| DL-histidine, HCl, H2O | Amino-acid | DHI; HIS | 24; n/a |
| DL-5-hydroxylysine, HCl | Amino-acid | n/a; LYZ | 0; 7 |
| Trans-4-hydroxy-L-proline | Amino-acid | HYP | 149 |
| Spermine, 4HCl | Polyamine | SPM | 103 |
| Spermidine, 3 HCl | Polyamine | SPD | 32 |
| 1,4-diaminobutane, 2 HCl | Polyamine | PUT | 22 |
| DL-ornithine, HCl | Polyamine | ORD; ORN | 3; 56 |
| NDSB 256 | Surfactant | DMX | 4 |
| NDSB 195 | Surfactant | NDS | 7 |
| Bis-tris | Buffer | BTB | 114 |

*No of structures as determined by a query of the pdb carried out in December 2014

The Morpheus® Additive Screen MD1-93 Wells A1-D12

| Well \# | Chemical name | known as/for | Conc |  |
| :--- | :--- | :--- | :--- | :--- |
| A1 | Water | control |  | - |
| A2 | PEG 20000 | precipitant | 20.00 | $\%$ |
| W/v |  |  |  |  |
| A3 | PEG 500 MME | precipitant | 40.00 | $\%$ |

The Morpheus@ Additive Screen MD1-93 Wells E1-H12

| Well \# | Chemical name | known as/for | Conc |  |
| :--- | :--- | :--- | :--- | :--- |
| E1 | Glycine | stabilizer | 0.40 | M |
| E2 | DL-Serine | stabilizer | 0.40 | M |
| E3 | PEG 3000 | precipitant | 30.00 | $\%$ |
| E4 | 1,2,4-Butanetriol | cryoprotectant | 40.00 | $\%$ |

${ }^{\dagger}$ A light pellet may form in tubes containing Lanthanides (G7-G10). It is easily re-suspended with further gentle mixing.


[^0]:    Please Note
    The screen should preferably be frozen upon arrival and defrosted prior to use
    Gently mix the screen before use (invert the block several times then spin down <1000 rpm).
    A light pellet may form in tubes containing Lanthanides (G7-G10). It is easily re-suspended with further gentle mixing.

