

| Stock | | Original call | Outcome date | Months ahead | Report title | Call | Outcome |
|--------|------|--------------------------|------------------|----------------|---|--|--|
| • | | | | Ave: 18 months | | | |
| CNTA | ALKS | 14/10/2025 | | | CNTA/ALKS/4502: CNTA faces orexin agonist hurdles | | |
| RARE | MREO | 15/09/2025 | | | RARE/MREO: ORBIT phase II: Improvements not solely attributable to natural progress seen with age | | |
| NOVN | OTSK | 14/05/2025 | Oct-25 | 5 | IGaN: VERA/OTSK/NOVN: oral Fablata'S likely eGFR benefit will leave injectable BAFF/APRIL's undifferentiated | APPLAUSE trial will show eGFR benefit | APPLAUSE P3 had eGFR benefit, leaving FDA approved Fabhalta as only oral |
| VERA | OTSK | 14/05/2025 | Jun-25 | 1 | IGaN: VERA/OTSK/NOVN: atacicept may lead IGaN | VERA ORIGIN3 P3 data will not be worse than competitors | ORIGIN3 well ahead of expectations |
| JNJ | | 08/05/2025 | | | JNJ: orexin antagonist seltorexant underappreciated in insomnia | | |
| JNJ | AZN | 08/05/2025 | | | JNJ: 1L Lazcluze + Rybrevant not convincing enough to topple Tagrisso | | |
| JNJ | | 08/05/2025 | | | JNJ: DUET 4804 IL23+antiTNF combo concerns for long-term tolerability | | |
| JNJ | ARGX | 09/05/2025 | | | JNJ/ARGX/UCB: FcRn Imaavy may struggle in MG, but is unique in Hemolytic disease of the newborn | | |
| SNYC | | 09/05/2025 | | | RPGR: All hope not lost for Beacon (SYNC) | | |
| PTGX | JNJ | 10/03/2025 | | | PTGX/JNJ:Icotrokinra UC efficacy undifferentiated to FDA approved oral competitors | | |
| NOVN | ALPS | 13/11/2024 | | | NOVN/APLS: Fabhalta Oral route opens the possibility of preventing AMD | | |
| AZN | NOVN | 13/11/2024 | | | AZN/NOVN: Myasthenia Refractory population and only partial inhibition of complement leaves Fabhalta likely in third place | | |
| NOVN | AZN | 13/11/2024 | | | NOVN/AZN:Atypical Haemolytic Uraemic syndrome: Fabahlta oral formulation differentiation should be a winner | | |
| NOVN | | 13/11/2024 | | | NOVN/AMGN: pelacarsen may be the first cardiovascular breakthrough in decades by lowering unconventional Lp(a) cholesterol to prevent heart attacks | | |
| ALPS | NOVN | 13/11/2024 | | | APLS/NOVN:C3G Fabhalta playing second fiddle to Empaveli | | |
| NOVN | | 13/11/2024 | Sep-25 | 10 | NOVN: Remibrutinib's clear path to \$4bn monopoly in Urticaria with a better BTK | Remibrutinib will be FDA approved | Remibrutinib FDA approved |
| NOVN | INI | 13/11/2024 | Aug-25 | 9 | NOVN/JNJ/AMGN: Ianalumab distinctive Sjögren's syndrome data, now first mover | Successful phase III NEPTUNUS . Will be 1st mkt. No PRO | Successful phase III NEPTUNUS .1st mkt. No PRO |
| SMMT | | 23/09/2024 | 7146 20 | Ŭ | SMMT: Hold your HARMONIs: VEGF in Asian lung cancer always did better | ouccoolar phase in the testing of the second the | |
| ROG | NOVN | 21/08/2024 | | | ROG/NOVN: Factor B lacking the X Factor in IGaN | | |
| NOVN | | 29/07/2024 | Jun-25 | 10 | NOVN/ROG/APLS: Fabulous Fabhalta in PNH halts Piasky blue skies | Improve switching +ve APPULSE | APPULSE +ve switch from C5 inc Hb by 2 |
| ROG | MRK | | Juli-25 | 10 | ROG/MRK/PFE: Roche's RVT3101 similar to incumbents in Ulcerative Colitis | Improve switching ive Air OLOL | AFFOLSE THE SWILLTHOM COME THO BY 2 |
| SRPT | ROG | 09/07/2024 29/05/2024 | Jun-24 | 1 | ROG/SRPT: Duchenne EMBARK likely enough to lessen FDA age restrictions | FDA full approval and no age restriction | Wide label FDA approval of Duchenne Gene Rx June 24 |
| | | | | 2 | CYTK: Aficamten approvable, may not outshine BMY's Camzyos | | |
| CYTK | BMY | 29/02/2024 | May-24 | | | Aficapmten underlying data not better than Camzyos | Take out premium lost after royalty deal |
| MRK | | 15/01/2024 | Mar-24 | 2 | MRK: Pul HT: High chance FDA, major share of high-priced orphan market; trial in heart failure could open huge market | FDA approval in Group 1. Chance in lareg Group 2 market | FDA approval of Pulmonary HT drug |
| MRK | | 15/01/2024 | | | MRK: TL1A: PRA023 despite 1st in class, may struggle to take significant share in ulcerative colitis, based on current phase II data | O CLU WOODAL LIDIDOCCI | O CALL WOODH INDOVIA |
| MRK | AMGN | 15/01/2024 | Nov-25 | 22 | MRK:MK0616 oral PCSK9 headline benefit came in undertreated patients, suggesting it may struggle to break into the large, now generic primary prevention market | Successful phase III CORAL-LIPIDS trial | Successful phase III CORAL-LIPIDS trial |
| MRK | AZN | 15/01/2024 | | | MRK: crashing the ADC party late with an attempt at differentiation that may not better competitors | | |
| MRK | | 15/01/2024 | Nov-25 | 22 | MRK: Sotatercept: Pulmonary HT CADENCE-PH trial in heart failure could open huge market | PH grooup 2 likley to be successful phase 2 | Phase 2 met primary endpoint of reduced PVR |
| BMY | BAY | 21/11/2023 | Nov-25 | 24 | BMY/JNJ/BAYN: Bayer AF failure does not reduce chances of success for ischaemic stroke trials for BMY/JNJ | OCEANIC-STROKE asundexian will redice stroke | OCEANIC-STROKE positive |
| AZN | GILD | 31/10/2023 | Sep-24 | 10 | AZN/GILD/Daiicii- ADCs in Lung cancer face steep hurdles after ESMO23 | Dato will miss OS in 2L and in 1L NSCLC | Dato-DXD misses OS in Lung and Breast |
| PTGX | JNJ | 14/08/2023 | Mar-25 | 19 | PTGX/JNJ: Oral IL23 dosing faces hurdles of new rival oral classes and psoriasis in ulcerative colitis | | |
| MRNA | MRK | 19/04/2023 | | | MRK/MRNA: mRNA-PD1 cancer vaccine trial control arm may have underperformed | | |
| MRK | | 20/03/2023 | Dec-23 | 9 | MRK: Little hope in TIGIT trials to date | TIGIT class will fail | TIGIT fails |
| Pfizer | SAN | 21/12/2022 | Oct-24 | 22 | PFE: pregnancy RSV vaccine: \$450m. Stiff competition from single-use antibody targeted to high-risk children limits utility. | RSV vaccine will not take share from Beyfortus | Beyfortus antibody continues to grow 382% a year later |
| Pfizer | | 21/12/2022 | Oct-23 | 9 | PFE: 2024: Etrasimod: Ulcerative colitis: Good chance of FDA approval, better practical oral efficacy in large unmet market | Etrasimod FDA approval in UC | Fda approval of Velsipity in UC |
| Pfizer | | 21/12/2022 | Aug-23 | 7 | PFE: Elranatamab: Myeloma: faces stiff competition from ever rising CAR-Ts | Elranatamab undifferentiated | 1 year after launch, 2024E only \$100m |
| Pfizer | | 21/12/2022 | Jun-23 | 5 | PFE: Ritlecitinib: Alopecia areata: 2023: High chance of FDA, majority share in large, high priced unmet need, despite 2nd to market with better efficacy | Ritlecitinib FDA approval | FDA approval of Litfulo in Hair Loss |
| CNCE | | 21/12/2022 | Jan-23 | 1 | CNCE: deuruxolitinib appear gain a response in more patients, and faster than baricitinib | Deuruxolitinib appears superior to baricitinib | Alopeica BuyOut by Sun |
| ROG | | 15/08/2022 | Feb-23 | 6 | ROG: Idiopathic pulmonary fibrosis: PRM-151 tepid | PRM-151 undirrefentiete to standard of care, will likley fail | STARSCAPE terminated for futility |
| ROG | AZN | 15/08/2022 | Oct-25 | 38 | ROG: Giredestrant needs restriction to very narrow ESR1 or will likely fail breast Ca | PFS will be driven by ESR1 benefit only | PFS was driven by ESR1 benefit only |
| BMY | | 21/06/2022 | Jun-22 | 1 | BMY/Kite: Bristol's Breyanzi best in class | FDA approval in LBCL | Breyanzi FDA approved |
| BAY | BMY | 16/05/2022 | Nov-23 | 18 | BAY: Bayer's Factor Xia lower bleeding came with more stroke in phase II AF | Asundexian unlikely to succeed in AF or MI | Asundexian fails in AF or heart attacks |
| JNJ | | 08/03/2022 | Oct-24 | 31 | JNJ: TAR200 MIBC: High 50% repsonse rates, though only very early data | TAR-200 success in adj MIBC | MIBC: fails adj phase III vs chemoradiation |
| JNJ | | 08/03/2022 | Sep-24 | 30 | JNJ: TAR200 MIBC: High 50% repsonse rates, though only very early data | TAR-200 success in neo adj MIBC | MIBC: positive neo adj phase II vs PD1 |
| JNJ | | 08/03/2022 | Jun-24 | 27 | JNJ: Nipocalimab: Likley successful phase III, less shots, longer duration, but late to market | Nipocalimab postive phase III in MG | Positive phase III MG trial |
| JNJ | | 08/03/2022 | Apr-23 | 13 | JNJ: TAR-200 likley FDA approval in NMIBC | FDA approval in NMIBC | Positive nBCG-NMIBC registrational study: FDA filed |
| JNJ | | 08/03/2022 | Oct-22 | 7 | JNJ: Teclistamab phase II similar efficacy but no G3 neurotoxicity may be competitive threat to Carvykti | FDA approval with less neurotox than Carvykti | FDA approved, but Black box neurotoxic in phase III: Black box: but less than Carvkyti |
| GILD | | 10/02/2022 | Feb-23 | 12 | GILD: Trodelcy concern: Approveable, but high risk to be able to better chemo in previously CDK treated patients | FDA approval, but no benefit in prior CDK | FDA approves Trodelvy in HER2-ER+ 2L BC, with no effect in fully CDK treated pts |
| BMY | | 11/07/2021 | Oct-24 | 39 | BMY:Cendakimab: eosinophilic oesophagitis likely successful in steroid refractory | Successful Phase III | Successful Phase III |
| ВМҮ | | 11/07/2021 | Sep-22 | 14 | BMY: Deucravacitinib: plaque psoriasis: 2022: likely FDA approval, better than Otezla, safer than JAKs, easier to take than Stelara | FDA approval of Sotyktu in psoriasis | FDA approval of Sotyktu in psoriasis |
| ВМҮ | | 11/07/2021 | Aug-22 | 13 | BMY: Mavacamten: cardiomyopathy: 2022:FDA approval very likely. | FDA approval of Camzyos in HoCM | FDA approval of Camzyos in HoCM |
| BAY | BAY | 11/07/2021 | Nov-25 | 53 | Factor XIa inhibitor BMS-986177 ischaemic stroke prevention 2025: early data and epidemiology suggest new gold standard | OCEANIC-STROKE asundexian will redice stroke | OCEANIC-STROKE positive |
| ВМҮ | | 28/06/2021 | Mar-22 | 8 | BMY: LAG3+Opdivo approvable, but driven by PD1-ve only | FDA approval of Opdulag | FDA appoved Opdulag |
| GSK | AZN | 02/02/2021 | Feb-23 | 24 | GSK: Daprodustat FDA approval likely only in dialysis dependent. | Restricted FDA approval to dialysis-dependent only | FDA Daprodustat approval restricted to dialysis dependent only |
| JNJ | | 02/02/2021 | | | Zejula in lung doubtful | , , , , , , , , , , , , , , , , , , , | · · · · · · · · · · · · · · · · · · · |
| JNJ | | 02/02/2021 | Feb-24 | 36 | Gepotidacin good prospects of approval, though needs time for standard of care to become ineffective | Successful phase III, FDA approval | Positive phase III EAGE trial.FDA filed |
| AZN | | 02/02/2021 | Dec-21 | 10 | cabotegravir prep likely FDA approval, but faces 1st gen generics | FDA approval | Cabotegravir FDA approved |
| JNJ | | 15/10/2020 | Apr-22 | 18 | Mayacamten's potential | Successful phase III, FDA approval | FDA appoved mavacempten |
| ROG | INII | 23/09/2020 | Aug-24 | 47 | JNJ: amivantamab + lazertinib combo data is supportive of first line success | Will beat Tagrisso in 1L; FDA approval | FDA approved lazertinib+ amivantamab in 1st line EGFR NSCLC with exon 19/21 |
| JNJ | AZN | 23/09/2020 | Aug-23 | 35 | JNJ: Niraparib: likely FDA (MAGNITUDE) in 1L prostate cancer, but late and undifferentiated | FDA approval, but undifferentiated to Lynparza | FDA approval of Niraparib in 1st line suspected or deleterious BRCA prostate cancer |
| JNJ | AZN | | | 32 | AZN: Lynparza PROpel will likely lead to 1st line PFS in ITT popn | Wider FDA approval | FDA approved Lynparza in 1st line suspected or deleterious BRCA prostate cancer |
| JNJ | AZN | 23/09/2020 23/09/2020 | May-23 May-21 | 32 8 | JNJ: Amivantamab good chance of FDA success in 1st line exon 20 insertion lung cancer | FDA apporval in ex20i | Amivantamab FDA-approved Exon20i |
| ROG | ALIV | | | | | RocheTIGIT will fail | Roche TIGIT fails |
| | | 12/06/2020 | Apr-22 | 22 | ROG.SW: TIGIT tweaks trailing Tecentriq but faces far more effective rivals Tak721: Likely Eta approval in ensingnibilic geson haritis | | |
| Takeda | | 19/04/2020 | Feb-24 | 46 | Tak721: Likely Fda approval in eosinophilic oesophagitis TAK 799: Specific mutated "Llung capeer likely EDA approval in 2021 | FDA approval | FDA approved on second filing |
| Takeda | | 19/04/2020 | Sep-21 | 17 | TAK-788: Specific mutated "I lung cancer, likely FDA approval in 2021 | FDA approval | FDA approved |
| Takeda | | 19/04/2020 | Nov-21 | 19 | TAK-620: post-transplant cytomegalovirus likely FDA approval in 2021 | FDA approval | FDA approved |
| Takeda | | 19/04/2020 | Jul-22 | 27 | TAK-924: Concerns over response rate data will deliver OS benefit | Phase III likley to fail | TAK-924 fails phase III |
| Takeda | | 19/04/2020 | May-22 | 25 | TAK-609: intrathecal Elaprase in Hunter's unlikely to get FDA approval | FDA failure | TAK-609 discontinued |
| NOVN | | 23/02/2020 | Jun-23 | 40 | NOVN: Kisqali's Kiss of life: Unique survival and tolerability bodes well for adjuvant | Successful phase III in adj, wider use than Verzenio; FDA adjuvant | |
| NOVN | | 23/02/2020 | Aug-20 | 5 | NOVN: Ofatumumab: best MS efficacy, Likley Fda approval, £3bn 2024 conservative estimate | | FDA approves Ofatumumab (2024 sales ~\$3.6bn |
| NOVN | AMGN | 23/02/2020 | Dec-21 | 22 | NOVN: inclisiran likely to get FDA approval this year;looks likely to easily become blockbuster | | FDA approves inclisiran. Current \$1bn annual run rate |
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