**Supplementary Figures**

**Figure 1**

**Fig S1. Acquired drug resistance enhances migration and invasion of CRC cells.** RKO cells were cultured in 3D Matrigel with 10 µM 5'-FU and 15 µM CB-839 for 4 weeks to select for drug-resistant clones. One of the resistant clones was retested for resistance by another round of treatment on equal number of RKO and RKO-DR cells with both the drugs. After 9 days, cytotoxicity was determined by measuring cell counts. Bars indicate mean ± sem. *p<0.0001 by unpaired t-test. Increased invasion through Matrigel layer and migration of the non-resistant and drug-resistant CRC cell lines was measured using transwell assays. Cells were stained using 0.1% crystal violet. After drying, the wells were imaged using Moticam T2 camera, and the cells were counted using FIJI software. Compared to their parent counterparts, the drug-resistant B. DLD1-DR and C. RKO-DR lines showed significantly higher number of cells invading and migrating through the transwells. Bars indicate mean ± sem. *p<0.05 using unpaired t-test.
Fig S2. Time course of contrast-enhanced MRMI of EDB-FN using MT218 facilitates non-invasive assessment of drug resistance in DLD-1 tumors. T1-weighted FSE coronal and axial images were obtained pre- and post-injection (at varying time-points) of 40 μmol/kg dose of MT218 in DLD-1 and DLD1-DR xenograft-bearing athymic nu/nu female mice. Representative A. coronal and B. axial images show steady increase in signal enhancement for up to 30 min in both DLD-1 and DLD1-DR tumors, with increased enhancement in the latter ones. Significant CNR enhancement is observed in C. coronal and D. axial images up to 25-30 min in both the tumors compared to pre-contrast, with significantly higher CNR in the DLD1-DR tumors, compared to DLD-1 tumors. (n=2 mice per group, 2 slices per mouse). Dots indicate mean ± sem. *p<0.05 (time course) and **p<0.005 (DLD-1 vs DLD1-DR) using 2-way ANOVA with Tukey’s correction.
Fig S3. Time course of contrast-enhanced MRMI of EDB-FN using MT218 facilitates non-invasive assessment of drug resistance in RKO tumors. T1-weighted FSE coronal and axial images were obtained pre- and post-injection (at varying time-points) of 40 µmol/kg dose of MT218 in RKO and RKO-DR xenograft-bearing athymic nu/nu female mice. Representative A. coronal and B. axial images show steady increase in signal enhancement for up to 30 min in both RKO and RKO-DR tumors, with increased enhancement in the latter ones. Significant CNR enhancement is observed in C. coronal and D. axial images up to 25-30 min in both the tumors compared to pre-contrast, with significantly higher CNR in the RKO-DR tumors, compared to RKO tumors. (n=2 mice per group, 2 slices per mouse). Dots indicate mean ± sem. ***p<0.0001 for time course and RKO vs RKO-DR for C and **p<0.01 for time course and RKO vs RKO-DR for D, using 2-way ANOVA with Tukey’s correction.
Fig S4. MRMI of EDB-FN with MT218 for non-invasive therapeutic monitoring of mice bearing drug-resistant CRC tumors. CNRs of T₁-weighted 2D spin echo axial images were obtained from MRMI performed at Baseline (week 1) and Endpoint (week 4) with 40 µmol/kg dose MT218 in DLD1-DR-bearing mice treated with DMSO (vehicle group) and MK2206-HCl (treated group). No significant difference observed in the average CNRs between both the groups at week 1. While the week 4 average CNR of the treated group is higher than that of the vehicle group, it is not statistically significant. Bars indicate mean ± sem and dots indicate individual mouse tumor volumes. n=5 mice per group, *p<0.05 using unpaired t-test.